

Europäisches Patentamt European Patent Office

Office européen des brevets



(11) EP 0 979 281 B1

(12)

EUROPEAN PATENT SPECIFICATION

- (45) Date of publication and mention of the grant of the patent:

 20.07.2005 Bulletin 2005/29
- (21) Application number: 98920059.7
- (22) Date of filing: 30.04.1998

- (51) Int CI.7: **C12N 15/13**, C07K 16/46, C12N 5/10, C07K 16/28, C07K 16/32
- (86) International application number: PCT/US1998/008762
- (87) International publication number: WO 1998/050431 (12.11.1998 Gazette 1998/45)

(54) A METHOD FOR MAKING MULTISPECIFIC ANTIBODIES HAVING HETEROMULTIMERIC AND COMMON COMPONENTS

ein verfahren zur herstellung multispezifischer antikörper die heteromultimere und gemeinsame komponenten besitzen

PROCEDE DE PREPARATION D'ANTICORPS MULTISPECIFIQUES PRESENTANT DES COMPOSANTS HETEROMULTIMERES

- (84) Designated Contracting States:

 AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU

 MC NL PT SE
- (30) Priority: **02.05.1997 US 850058 24.06.1997 US 50661 P**
- (43) Date of publication of application: **16.02.2000 Bulletin 2000/07**
- (73) Proprietor: Genentech, Inc.
 South San Francisco, CA 94080-4990 (US)
- (72) Inventors:
 - ARATHOON, Robert San Mateo, CA 94402 (US)
 - CARTER, Paul, J. San Francisco, CA 94127 (US)
 - MERCHANT, Anne, M.
 San Bruno, CA 94116 (US)
 - PRESTA, Leonard, G.
 San Francisco, CA 94109 (US)
- (74) Representative: Walton, Seán Malcolm et al Mewburn Ellis LLP
 York House,
 23 Kingsway
 London WC2B 6HP (GB)
- (56) References cited:

WO-A-92/10209

WO-A-96/27011

- VAUGHAN ET AL: "HUMAN ANTIBODIES WITH SUB-NANOMOLAR AFFINITIES ISOLATED FROM A LARGE NON-IMMUNIZED PHAGE DISPLAY LIBRARY" NATURE BIOTECHNOLOGY, vol. 14, 1996, pages 309-314, XP002084763 cited in the application
- ZHU ET AL: "REMODELLING DOMAIN INTERFACES TO ENHANCE HETERODIMER FORMATION" PROTEIN SCIENCE, vol. 6, no. 4, April 1997, pages 781-788, XP002084764
- WELLS ET AL: "IN VIVO FORMATION AND STABILITY OF ENGINEERED DISULFIDE BONDS IN SUBTILISIN" THE JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 261, no. 14, 15 May 1986, pages 6564-6570, XP002084765
- RIDGWAY ET AL: "'KNOBS-INTO-HOLES' ENGINEERING OF ANTIBODY CH3 DOMAINS FOR HEAVY CHAIN HETERODIMERIZATION" PROTEIN ENGINERING, vol. 9, 1996, pages 617-621, XP002084766 cited in the application
- FIGINI ET AL: "IN VITRO ASSEMBLY OF REPERTOIRES OF ANTIBODY CHAINS ON THE SURFACE OF PHAGE BY RENATURATION" JOURNAL OF MOLECULAR BIOLOGY, vol. 239, 1994, pages 68-78, XP002084767 cited in the application
- NISSIM ET AL: "ANTIBODY FRAGMENTS FROM A 'SINGLE POT' PHAGE DISPLAY LIBRARY AS IMMUNOCHEMICAL REAGENTS" THE EMBO JOURNAL, vol. 13, no. 3, 1994, pages 692-698, XP002084768

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

Description

5

10

20

25

40

45

50

55

FIELD OF THE INVENTION

[0001] This invention relates to a method for making multispecific antibodies having heteromultimericheavy chain components and common light chain components such as bispecific antibodies, bispecific immunoadhesins, as well as antibody-immunoadhesinchimeras and the heteromultimeric polypeptides made using the method.

BACKGROUND OF THE INVENTION

Bispecific antibodies

[0002] Bispecific antibodies (BsAbs) which have binding specificities for at least two different antigens have significant potential in a wide range of clinical applications as targeting agents for *in vitro* and *in vivo* immunodiagnosis and therapy, and for diagnostic immunoassays.

[0003] In the diagnostic areas, bispecific antibodies have been very useful in probing the functional properties of cell surface molecules and in defining the ability of the different Fc receptors to mediate cytotoxicity(Fanger *et al.*, Crit. Rev. Immunol. 12:101-124 (1992)). Nolan *et al.*, Biochem. Biophys. Acta. 1040:1-11 (1990) describe other diagnostic applications for BsAbs. In particular, BsAbs can be constructed to immobilize enzymes for use in enzyme immunoassays. To achieve this, one arm of the BsAb can be designed to bind to a specific epitope on the enzyme so that binding does not cause enzyme inhibition, the other arm of the BsAb binds to the immobilizing matrix ensuring a high enzyme density at the desired site. Examples of such diagnostic BsAbs include the rabbit anti-IgG/anti-ferritin BsAb described by Hammerling *et al.*, J. Exp. Med. 128:1461-1473 (1968) which was used to locate surface antigens. BsAbs having binding specificities for horse radish peroxidase (HRP) as well as a hormone have also been developed. Another potential immunochemical application for BsAbs involves their use in two-site immunoassays. For example, two BsAbs are produced binding to two separate epitopes on the analyte protein - onc BsAb binds the complex to an insoluble matrix, the other binds an indicator enzyme (see Nolan *et al.*, *supra*).

Bispecific antibodies can also be used for in vitro or in vivo immunodiagnosis of various diseases such as cancer (Songsivilaiet al., Clin. Exp. Immunol. 79:315 (1990)). To facilitate this diagnostic use of the BsAb, one arm of the BsAb can bind a tumor associated antigen and the other arm can bind a detectable marker such as a chelator which tightly binds a radionuclide. Using this approach, Le Doussal et al. made a BsAb useful for radioimmunodetectionof colorectal and thryoid carcinomas which had one arm which bound a carcinoembryonicantigen (CEA) and another arm which bound diethylenetriaminepentacetic acid (DPTA). See Le Doussal et al., Int. J. Cancer Suppl. 7:58-62 (1992) and Le Doussal el al., J. Nucl. Med. 34:1662-1671 (1993). Stickney et al. similarly describe a strategy for detecting colorectal cancers expressing CEA using radioimmunodetection. These investigators describe a BsAb which binds CE_A as well as hydroxyethylthiourea-benryl-EDTA (EOTUBE). See Stickney et al., Cancer Res. 51:6650-6655 (1991). [0005] Bispecific antibodies can also be used for human therapy in redirected cytotoxicity by providing one arm which binds a target (e.g. pathogen or tumor cell) and another arm which binds a cytotoxic trigger molecule, such as the Tcell receptor or the Fcγ receptor. Accordingly, bispecific antibodies can be used to direct a patient's cellular immune defense mechanisms specifically to the tumor cell or infectious agent. Using this strategy, it has been demonstrated that bispecific antibodies which bind to the FcγRIII (i.e. CD16) can mediate tumor cell killing by natural killer (NK) cell/ large granular lymphocyte (LGL) cells in vitro and are effective in preventing tumor growth in vivo. Segal et al., Chem. Immunol. 47:179 (1989) and Segal et al., Biologic Therapy of Cancer 2(4). DeVita et al. eds. J.B. Lippincott, Philadelphia (1992) p. I. Similarly, a bispecificantibodyhaving one arm which binds FcγRIII and another which binds to the HER2 receptor has been developed for therapy of ovarian and breast tumors that overexpress the HER2 antigen. (Hseih-Ma et al. Cancer Research 52:6832-6839 (1992) and Weiner et al. Cancer Research 53:94-100 (1993)). Bispecific antibodiescan also mediate killing by T cells. Normally, the bispecific antibodies link the CD3 complex on T cells to a tumorassociatedantigen. A fully humanized F(ab')₂ BsAb consisting of anti-CD3 linked to anti-p185^{HER2} has been used to target T cells to kill tumor cells overexpressingthe HER2 receptor. Shalaby et al., J. Exp. Med. 175(1):217 (1992). Bispecific antibodies have been tested in several early phase clinical trials with encouraging results. In one trial, 12 patients with lung, ovarian or breast cancer were treated with infusions of activated T-lymphocytes targeted with an anti-CD3/anti-tumor (MOC31) bispecific antibody. deLeij et al. Bispecific Antibodies and Targeted Cellular Cytotoxicity, Romet-Lemonne, Fanger and Segal Eds., Lienhart (1991) p. 249. The targeted cells induced considerable local lysis of tumor cells, a mild inflammatory reaction, but no toxic side effects or anti-mouse antibody responses. In a very preliminary trial of an anti-CD3/anti-CD 19 bispecific antibody in a patient with B-cell malignancy, significant reduction in peripheral tumor cell counts was also achieved. Clark et al. Bispecific Antibodies and Targeted Cellular Cytotoxicity, Romet-Lemonne, Fanger and Segal Eds., Lienhart (1991) p. 243. See also Kroesen et al., Cancer Immunol. Immunother. 37:400-407 (1993), Kroesen et al., Br. J. Cancer 70:652-661 (1994) and Weiner et al., J. Immunol. 152:2385

(1994) concerning therapeutic applications for BsAbs.

5

20

25

30

40

45

50

55

[0006] Bispecific antibodiesmay also be used as fibrinolyticagents or vaccine adjuvants. Furthermore, these antibodies may be used in the treatment of infectious diseases (e.g. for targeting of effector cells to virally infected cells such as HIV or influenza virus or protozoa such as *Toxoplasma gondii*), used to deliver immunotoxinsto tumorcells, or target immune complexes to cell surface receptors (see *Fanger et al.*, *supra*).

[0007] Use of BsAbs has been effectively hindered by the difficulty of obtaining BsAbs in sufficient quantity and purity. Traditionally, bispecific antibodies were made using hybrid-hybridomatechnology (Millstein and Cuello, Nature 305: 537-539 (1983)). Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of 10 different antibody molecules, of which only one has the correct bispecific structure (see Fig. 1A). The purification of the correct molecule, which is usually done by affinity chromatography steps, is rather cumbersome, and the product yields are low. See, for example, (Smith, W., et al. (1992) Hybridoma 4:87-98; and Massimo, Y.S., et al. (1997) J. Immunol. Methods 201:57-66). Accordingly, techniques for the production of greater yields of BsAb have been developed. To achieve chemical coupling of antibody fragments, Brennan el al., Science 229:81 (1985) describe a procedure wherein intact antibodies are proteotyticattycteaved to generate F(ab')₂ fragments. These fragments are reduced in the presence of the dithiol complexing agent sodium arsenite to stabilize vicinal dithiols and prevent intermolecular disulfide formation. The Fab' fragments generated are then converted to thionitrobenzoate (TNB) derivatives. One of the Fab'-TNB derivatives is then reconverted to the Fab'-thiol by reduction with mercaptoethylamine and is mixed with an equimolar amount of the other Fab'-TNB derivative form the BsAb. The BsAbs produced can be used as agents for the selective immobilization of enzymes.

[0008] Recent progress has facilitated the direct recovery of Fab'-SH fragments from *E. coli*. which can be chemically coupled to form bispecific antibodies. Shalaby *et al.*, J. Exp. Med. 175:217-225 (1992) describe the production of a fully humanized BsAb F(ab')₂ molecule having one arm which binds p185^{HER2} and another arm which binds CD3. Each Fab' fragment was separately secreted from *E. coli*. and subjected to directed chemical coupling *in vitro* to form the BsAb. The BsAb thus formed was able to bind to cells overexpressing the HER2 receptor and normal human T cells, as well as trigger the lytic activity of human cytotoxic lymphocytesagainst human breast tumor targets. See also Rodrigues *et al.*, Int. J. Cancers (Suppl.) 7:45-50 (1992).

[0009] Various techniques for making and isolating BsAb fragments directly from recombinant cell cultures have also been described. For example, bispecific F(ab')₂ heterodimers have been produced using leucine zippers (Kostelny et al., J. Immunol. 148(5):1547-1553 (1992)). The leucine zipper peptides from the Fos and Jun proteins were linked to the Fab' portions of anti-CD3 and anti-interleukin-2 receptor (IL-2R) antibodies by gene fusion. The antibody homodimers were reduced at the hinge region to form monomers and then reoxidized to form the antibody heterodimers. The BsAbs were found to be highly effective in recruiting cytotoxic T cells to lyse HuT-102 cells in vitro. The advent of the "diabody" technology described by Hollinger et al., PNAS (USA) 90:6444-6448 (1993) has provided an alternative mechanism for making BsAb fragments. The fragments comprise a heavy chain variable domain (VH) connected to a light chain variable domain (V_L) by a linker which is too short to allow pairing between the two domains on the same chain. Accordingly, the V_H and V_L domains of one fragment are forced to pair with the complementary V_I and V_H domains of another fragment, thereby forming two antigen-binding sites. Another strategy for making BsAb fragments by the use of single chain Fv (sFv) dimers has also been reported. See Gruber et al. J. Immunol. 152: 5368 (1994). These researchers designed an antibody which comprised the V_H and V_L domains of an antibody directed against the T cell receptor joined by a 25 amino acid residue linker to the V_H and V_I domains of an anti-fluoresceinantibody. The refolded molecule bound to fluorescein and the T cell receptor and redirected the lysis of human tumor cells that had fluorescein covalently linked to their surface.

[0010] It is apparent that several techniques for making bispecific antibody fragments which can be recovered directly from recombinant cell culture have been reported. However, full length BsAbs may be preferable to BsAb fragments for many clinical applications because of their likely longer serum half-life and possible effector functions.

<u>Immunoadhesins</u>

[0011] Immunoadhesins(la's) are antibody-like molecules which combine the binding domain of a protein such as a cell-surface receptor or a ligand (an "adhesin") with the effector functions of an immunoglobulin constant domain. Immunoadhesins can possess many of the valuable chemical and biological properties of human antibodies. Since immunoadhesinscan be constructed from a human protein sequence with a desired specificity linked to an appropriate human immunoglobulin hinge and constant domain (Fc) sequence, the binding specificity of interest can be achieved using entirely human components. Such immunoadhesins are minimally immunogenic to the patient, and are safe for chronic or repeated use.

[0012] Immunoadhesins reported in the literature include fusions of the T cell receptor (Gascoigne *et al.*, Proc. Natl. Acad. Sci. USA <u>84</u>:2936-2940(1987)); CD4 (Capon *et al.*, Nature <u>337</u>:525-531 (1989); Traunecker *et al.*, Nature <u>339</u>: 68-70 (1989); Zettmeissl *et al.*, DNA Cell Biol. USA <u>9</u>:347-353 (1990); and Byrn *et al.*, Nature <u>344</u>:667-670 (1990)); L-

selectin or homing receptor (Watson *et al.*, J. Cell. Biol. 110:2221-2229 (1990); and Watson *et al.*, Nature 349:164-167 (1991)); CD44 (Aruffo *et al.*, Cell 61:1303-1313 (1990)); CD28 and B7 (Linsley *et al.*, J: Exp. Med. 173:721-730(1991)); CTLA-4 (Lisley *et al.*, J. Exp. Med. 174:561-569 (1991)); CD22 (Stamenkovic *et al.*, Cell 66:1133-1144 (1991)); TNF receptor (Ashkenazi *et al.*, Proc. Natl. Acad. Sci. USA 88:10535-10539 (1991); Lesslauer *et al.*, Eur. J. Immunol. 27: 2883-2886 (1991); and Peppel *et al.*, J. Exp. Med. 174:1483-1489 (1991)); NP receptors (Bennett *et al.*, J. Biol. Chem. 266:23060-23067 (1991)); inteferon γ receptor (Kurschner *et al.*, J. Biol. Chem. 267:9354-9360 (1992)); 4-1BB (Chalupny *et al.*, PNAS (USA) 89:10360-10364 (1992)) and IgE receptor α (Ridgway and Gorman, J. Cell. Biol. Vol. 115, Abstract No. 1448 (1991)).

[0013] Examples of immunoadhesins which have been described for therapeutic use include the CD4-lgG immunoadhesin for blocking the binding of HIV to cell-surface CD4. Data obtained from Phase I clinical trials in which CD4-lgG was administered to pregnant women just before delivery suggests that this immunoadhesin may be useful in the prevention of maternal-fetal transfer of HIV. Ashkenazi *et al.*, Intern. Rev. Immunol. 10:219-227 (1993). An immunoadhesin which binds tumor necrosis factor (TNF) has also been developed. TNF is a proinflammatory cytokine which has been shown to be a major mediator of septic shock. Based on a mouse model of septic shock, a TNF receptor immunoadhesin has shown promise as a candidate for clinical use in treating septic shock (Ashkenazi *et al.*, *supra*). Immunoadhesins also have non-therapeutic uses. For example, the L-selectin receptor immunoadhesin was used as an reagent for histochemical staining of peripheral lymph node high endothelial venules (HEV). This reagent was also used to isolate and characterize the L-selectin ligand (Ashkenazi *et al.*, *supra*). If the two arms of the immunoadhesin structure have different specificities, the immunoadhesin is called a "bispecific immunoadhesin" by analogy to bispecific antibodies. Dietsch *et al.*, J. Immunol. Methods 162:123 (1993) describe such a bispecific immunoadhesin combining the extracellular domains of the adhesion molecules, E-selectin and P-selectin. Binding studies indicated that the bispecific immunoadhesins from which it was derived.

25 Antibody-Immunoadhesin chimeras

5

10

15

20

30

40

45

50

55

[0014] Antibody-immunoadhesin (Ab/la) chimeras have also been described in the literature. These molecules combine the binding region of an immunoadhesin with the binding domain of an antibody.

[0015] Berg et al., PNAS (USA) 88:4723-4727 (1991) made a bispecific antibody-immunoadhesin chimera which was derived from murine CD4-IgG. These workers constructed a tetrameric molecule having two arms. One arm was composed of CD4 fused with an antibody heavy-chainconstant domain along with a CD4 fusion with an antibody light-chain constant domain. The other arm was composed of a complete heavy-chain of an anti-CD3 antibody along with a complete light-chain of the same antibody. By virtue of the CD4-IgG arm, this bispecific molecule binds to CD3 on the surface of cytotoxic T cells. The juxtaposition of the cytotoxic-cells and HIV-infected cells results in specific killing of the latter cells.

[0016] Wftile Berg *et al. supra* describe a bispecific molecule that was tetrameric in structure, it is possible to produce a trimeric hybrid molecule that contains only one CD4-IgG fusion. See Chamow *et al.*, J. Immunol. 153:4268 (1994). The first arm of this construct is formed by a humanized anti-CD3 κ light chain and a humanizedanti-CD3 γ heavy chain. The second arm is a CD4-IgG immunoadhesin which combines part of the extracellular domain of CD4 responsible for gp120 binding with the Fc domain of IgG. The resultant Ab/Ia chimera mediated killing of HIV-infected cells using either pure cytotoxic T cell preparations or whole peripheral blood lymphocyte (PBL) fractions that additionally included Fc receptor-bearing large granular lymphocyte effector cells.

[0017] In the manufacture of the multispecific antibody heteromultimers, it is desirable to increase the yields of the desired heteromultimer over the homomultimer(s). The current method of choice for obtaining Fc-containing BsAb remains the hybrid hybridoma, in which two antibodies are coexpressed (Milstein and Cuello, Nature 305:537-540 (1983)).

[0018] In hybrid hybridomas, heavy (H) chains typically form homodimers as well as the desired heterodimers. Additionally, light (L) chains frequently mispair with non-cognate heavy chains. Hence, coexpression of two antibodies may produce up to ten heavy and light chain pairings (Suresh, M.R., et al. Methods Enzymol. 121:210-228 (1986)). These unwanted chain pairings compromise the yield of the BsAb and inevitably impose significant, and sometimes insurmountable, purification challenges (Smith, et al. (1992) supra; and Massimo, et al. (1997) supra).

[0019] Antibody heavy chains have previously been engineered to drive heterodimerization by introducing sterically complementary mutations in multimerization domains at the C_H3 domain interface (Ridgway et al. Protein Eng. 9: 617-621 (1996)) and optimization by phage display as described herein. Chains containing the modified C_H3 domains yield up to approximately 90% heterodimer as judged by formation of an antibody/immunoadhesin hybrid (Ab/Ia). Heterodimerized heavy chains may still mispair with the non-cognate light chain, thus hampering recovery of the BsAb of interest.

SUMMARY OF THE INVENTION

10

15

20

25

30

40

45

50

55

[0020] This application describes a strategy which serves to enhance the formation of a desired heteromultimeric bispecific antibody from a mixture of monomers by engineering an interface between a first and second polypeptide for hetero-oligomerization by providing a common variable light chain to interact with each of the heteromeric variable heavy chain regions of the bispecific antibody. There are three possible hetero- and homomultimers that can form from a first and second polypeptide, each of which is, in turn, associated with a first and second light chain, respectively. This gives rise to a total of ten possible chain pairings (Fig. 1A). A method of enhancing the formation of the desired heteromultimer can greatly enhance the yield over undesired heteromultimers and homomultimers.

[0021] The preferred interface between a first and second polypeptide of the heteromultimeric antibody comprises at least a part of the C_H3 domain of an antibody constant domain. The domain of each of the first and second polypeptides that interacts at the interface is called the multimerization domain. Preferably, the multimerizationdomain promotes interaction between a specific first polypeptide and a second polypeptide, thereby increasing the yield of desired heteromultimer (Fig. 1B). Interaction may be promoted at the interface by the formation of protuberance-into-cavitycomplementary regions; the formation of non-naturally occurring disulfide bonds; leucine zipper; hydrophobic regions; and hydrophilic regions. "Protuberances" are constructed by replacing small amino acid side chains from the interface of the first polypeptide with larger side chains (e.g. tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the protuberances are optionally created on the interface of the second polypeptide by replacing large amino acid side chains with smaller ones (e.g. alanine or threonine). Where a suitably positioned and dimensioned protuberance or cavity exists at the interface of either the first or second polypeptide, it is only necessary to engineer a corresponding cavity or protuberance, respectively, at the adjacent interface. Non-naturally occurring disulfide bonds are constructed by replacing on the first polypeptide a naturally occurring amino acid with a free thiol-containing residue, such that the free thiol interacts with another free thiol-containingresidue on the second polypeptide such that a disulfide bond is formed between the first and second polypeptides (Fig. 1B).

[0022] Single chain Fv fragments from a large non-immunized phage display library (Vaughan, T.J. et al. (1996) Nature Biotechnology 14:309-314, herein incorporated by reference in its entirety) revealed V-gene usage in which V_H and V_L sequences derived from certain germline V-gene segments predominated, families predominated in the repertoire. Examples of chain promiscuity in the repertoire were noted in which a particular heavy or light chain is found in combination with different partner chains (Vaughan, T.J. et al. (1996) *supra*).

[0023] It is disclosed herein that the preparation of a desired heteromultimeric multispecific antibody is enhanced when a common light chain is provided to pair with each of the variable heavy chains of the multispecific antibody. Use of a common variable light chain reduces the number of monomers that must correctly pair to form the antigen binding domains by limiting the number of light chains from two or more light chains (in a bispecific or multispecificantibody, respectively, prior to disclosure of the instant invention) to one light chain (in a multispecific antibody of the invention, see Fig. 1C).

[0024] Accordingly, the invention relates to a method of preparing a heteromultimeric multispecific antibody, the antibody comprising 1) a first polypeptide and a second polypeptide (and additional polypeptides accord to the multiplicity of the antibody) which meet at an interface, wherein the first and additional polypeptides (*i.e.*, a first and second polypeptide) each include a multimerization domain forming an interface between the first and second (or at least one additional) polypeptides, and the multimerization domains promote stable interaction between first and additional polypeptides, and 2) a binding domain in each of the first and at least one additional polypeptide (*i.e.* a second polypeptide), each binding domain comprising a variable heavy chain and a variable light chain, wherein the variable light chain of the first polypeptide and the variable light chain of the second polypeptide have a common amino acid sequence, which common sequence has an amino acid sequence identity to an original light chain of each of the polypeptides of at least 80%, preferably at least 90%, more preferably at least 95% and most preferably 100% sequence identity. The method comprises the steps of

- (i) culturing a host cell comprising nucleic acid encoding the first polypeptide, the second polypeptide, and the common light chain wherein the culturing is such that the nucleic acid is expressed; and
- (ii) recovering the multispecific antibody from the host cell culture;

In a related embodiment of the invention the nucleic acid encoding the first polypeptide or the nucleic acid encoding the second polypeptide, or both, has been altered from the original nucleic acid to encode the interface or a portion thereof.

[0025] In another embodiment of the method, the interface of the first polypeptide comprises a free thiol-containing residue which is positioned to interact with a free thiol-containing residue of the interface of the second polypeptide such that a disulfide bond is formed between the first and second polypeptides. According to the invention, the nucleic acid encoding the first polypeptide has been altered from the original nucleic acid to encode the free thiol-containing

residue or the nucleic acid encoding the second polypeptide has been altered from the original nucleic acid to encode the free thiol-containing residue, or both.

[0026] In another embodiment of the method, the nucleic acid encoding both the first polypeptide and at least one additional polypeptide (*i.e.*, a second polypeptide) are altered to encode the protuberance and cavity, respectively. Preferably the first and second polypeptides each comprise an antibody constant domain such as the C_H3 domain of a human IgG₁.

[0027] In another aspect, the invention provides a heteromultimer (such as a bispecific antibody, bispecific immunoadhesin or antibody/immunoadhesinchimera) comprising a first polypeptide and a second polypeptide which meet at an interface. The interface of the first polypeptide comprises a multimerization domain which is positioned to interact with a multimerizationdomain on the at least one additional polypeptide (*i.e.*, a second polypeptide) to form an interface between the first and second polypeptide. In preferred embodiments of the invention, the multimerization domains are altered to promote interaction between a specific first polypeptide and a specific second polypeptide, which alterations include, but are not limited to, the generation of a protuberanceor cavity, or both; the generation of non-naturally occurring disulfide bonds; the generation of complementary hydrophobic regions; and the generation of complementary hydrophilic regions. The heteromultimeric multispecfic antibody may be provided in the form of a composition further comprising a pharmaceutically acceptable carrier.

[0028] The invention also relates to a host cell comprising nucleic acid encoding the heteromultimeric multispecificantibody of the preceding paragraph wherein the nucleic acid encoding the first polypeptide and at least one additional polypeptide (*i.e.*, a second polypeptide) is present in a single vector or in separate vectors. The host cell can be used in a method of making a heteromultimeric multispecific antibody which involves culturing the host cell so that the nucleic acid is expressed, and recovering the heteromultimeric antibody from the cell culture.

[0029] In yet a further aspect, the invention provides a method of preparing a heteromultimeric multispecific antibody comprising:

- (a) selecting a first nucleic acid encoding a first polypeptide comprising an amino acid residue in the interface of the first polypeptidethat is positioned to interact with an amino acid residue of interface of at least one additional polypeptide. In an embodiment the nucleic acid is altered from the original to encode the interacting amino acid residues. In another embodiment, the first nucleic acid is altered to encode an amino acid residue having a larger side chain volume, thereby generating a protuberance on the first polypeptide;
- (b) altering a second nucleic acid encoding a second polypeptide so that an amino acid residue in the interface of the second polypeptide is replaced with an amino acid residue having a smaller side chain volume, thereby generating a cavity in the second polypeptide, wherein the protuberance is positioned to interact with the cavity;
- (c) introducing into a host cell the first and second nucleic acids and culturing the host cell so that expression of the first and second nucleic acid occurs; and
- (d) recovering the heteromultimeric antibody formed from the cell culture.

[0030] It may also be desirable to construct a multispecific antibody (such as a bispecific antibody) that incorporates a previously identified antibody. Under these circumstances it is desirable to identify a heavy chain that when paired with the original light chain will bind specifically to a second antigen of interest. The methods of Figini *et al.* (Figini, M. *et al.* (1994) J. Mol. Biol. 239:68-78) may be used to identify such a heavy chain. First a phage library would be treated with guanidine hydrochloride to dissociate the original light chain. Next, the heavy chains displayed on phage would be reconstituted with the light chain of interest by removing the denaturant(such as by dialysis). Panning against the second antigen of interest would then be conducted to identify the desired heavy chain. The invention further embodies a multispecific antibody prepared by this method of selecting a heavy chain to pair with a chosen light chain, nucleic acid encoding the antibody, and a host cell comprising the nucleic acid.

[0031] The invention provides a mechanism for increasing the yields of the heteromultimerover other unwanted end-products such as undesired heteromultimers and/or homomultimers (see Fig. 1A-1C). Preferably, the yields of the desired heteromultimer recovered from recombinantcell culture are at least greater than 80% by weight and preferably greater than 90% by weight compared to the by-product undesired heterodimer or homomultimer(s).

Brief Description of the Drawings

[0032]

5

10

15

20

25

30

35

40

45

50

55

Figs. 1A-1C. Fig. 1A is a diagram of the formation of Fc-containing bispecific antibodies when no engineering is performed to enhance heteromuttimerizationover homomuttimerization. Fig. 1B is a diagram showing pairing that occurs when heavy (H) chains are engineered such that desired heteromultimerization is favored over undesired heteromultimerization over homomultimerization. Fig. 1C is a diagram showing pairing that occurs when antibodies

are chosen which share the same light (L) chain to circum vent the problem of light chains pairing with non-cognate heavy chains.

Figs. 2A-2C. Fig. 2A diagrams a selection scheme for C_H3 heterodimer using phage display vector, pRA2. Phage displaying stable C_H3 heterodimersare captured using an antibody directed to the gD flag. Fig. 2B diagrams a dicistronicoperon in which C_H3 expressed from a synthetic gene is co-secreted with a second copy of C_H3 expressed from the natural gene (Ellison et al. Nucleic Acids Res. 10:4071-4079 (1982)) as a fusion protein with M13 gene III protein. The synthetic C_H3 gene is preceded by a sequence encoding a peptide derived from herpes simplex virus glycoprotein D (gD flag, Lasky, L. A. and Dowbenko, D. J. (1984) DNA 3:23-29; Berman, P. W. et al., (1985) Science 227:1490-1492 and a cleavage (G) site for the site-specific protease, Genenase I (Carter, P. et al. (1989) Proteins: Structure, Function and Genetics 6:240-248). Fig. 2C is the nucleic acid sequence of the dicistronic operon (SEQ ID NO:1) of Fig. 2B in which the residues in the translated C_H3 genes are numbered according to the Eu system of Kabat et al. In Sequences of Proteins of Immunological Interest, 5th ed. vol. 1, pp. 688-696, NIH, Bethesda, MD (1991). Protuberance mutation T366W is shown, as are the residues targeted for randomization in the natural C_H3 gene (366, 368, and 407).

Figs. 3A-3C. Figs. 3A and 3B are bar graphs of the results of scanning densitometric analysis of SDS-PAGE of protein A-purified products from cotransfection of antibody (Ab) heavy and light chains with immunoadhesin(Ia). Data presented are the mean of two independent experiments. The x-axis indicates the ratios of input DNA by mass (Ia:H:L) and the y-axis indicates the percentage of each type of product multimer with respect to total product protein. Fig. 3C is a diagram of the possible product multimers.

Fig. 4 is a comparison of the V_L sequences of eight different antibodies with specificities for AxI, Rse, IgER, Ob-R, and VEGF. The position of the antigen binding CDR residues according to sequence definition (Kabat et al. (1991) *supra*) or structural definition (Chothia, C. and Lesk, A.M. J. Mol. Biol. (1987) 196:901-917) are shown by underlining and #, respectively. Residues that differ from the Ax1.78 sequence are shown by double underlining. Fig. 5 is a comparison of the heavy and light chains of selected anti-Ob-R and anti-HER3 clones. Shown are the V_H and the common V_L sequences of anti-Ob-R clone 26 and anti-HER3 clone 18 used to construct a bispecific antibody.

Fig. 6. Sandwich ELISA for detection of simultaneous binding to Mpl-IgG and HER3-IgG. Antibodies tested d were the anti-Mp1 x anti-HER3 BsIgG containing the mutations, Y349C:T366S:L368A:Y407V/T366'W:S354'C, together with corresponding parental anti-Mpl or anti-HER3 IgG with mutated Fc regions.

Fig. 7 is a bar graph of the results of an antibody-dependent cell-mediated cytotoxicity (ADCC) study. ADCC was mediated by huMAb4D5-5 (Carter, P. et al. (1992) PNAS USA 89:4285-4289) containing either a mutant (S354C: T366W/Y349'C:T366'S:L368'A:Y407'V) or wild-type Fc or an isotype-matched control antibody (E25, Presta, L.G. et al. (1993) J. Immunol. 151:2623-2632). The antibodies (125 ng/ml) were incubated with human peripheral blood mononuclear effector cells and SK-BR-3 target cells at the ratios shown. Data presented are the mean of triplicate measurements and representative of three separate experiments.

Fig. 8 is a matrix representing the amino acid sequence identity between the light chains of antibodies raised to HER3 versus the light chains of antibodies raised to Ob-R. Antibodies having light chains with 100% sequence identity are indicated in blackened boxes. Antibodies having light chains with 98-99% sequence identity are indicated in white boxes. The antibody clone identity is indicated below the matrix.

I. Definitions

5

10

15

20

25

30

35

40

45

50

55

[0033] In general, the following words or phrases have the indicated definitions when used in the description, examples, and claims:

[0034] A "heteromultimer", "heteromultimeric polypeptide", or "heteromultimeric multispecific antibody" is a molecule comprising at least a first polypeptide and a second polypeptide, wherein the second polypeptide differs in amino acid sequence from the first polypeptide by at least one amino acid residue. Preferably, the heteromultimerhas binding specificity for at least two different ligands or binding sites. The heteromultimer can comprise a "heterodimer" formed by the first and second polypeptide or can form higher order tertiary structures where polypeptides in addition to the first and second polypeptide are present. Exemplary structures for the heteromultimer include heterodimers (e.g. the bispecific immunoadhesin described by Dietsch et al., supra), heterotrimers (e.g. the Ab/la chimera described by Chamow et al., supra), heterotetramers (e.g. a bispecific antibody) and further oligomeric structures.

[0035] As used herein, "multimerization domain" refers to a region of each of the polypeptides of the heteromultimer. The "multimerizationdomain" promotes stable interaction of the chimeric molecules within the heteromultimer complex. Preferably, the multimerization domain promotes interaction between a specific first polypeptide and a specific second polypeptide, thereby enhancing the formation of the desired heteromultimer and substantially reducing the probability of the formation of undesired heteromultimers or homomultimers. The multimerizationdomains may interact via an immunoglobulin sequence, leucine zipper, a hydrophobic region, a hydrophilic region, or a free thiol which forms an

intermolecular disulfide bond between the chimeric molecules of the chimeric heteromultimer. The free thiol may be introduced into the interface of one or more interacting polypeptides by substituting a naturally occurring residue of the polypeptide with, for example, a cysteine at a position allowing for the formation of a disulfide bond between the polypeptides. The multimerizationdomain may comprise an immunoglobulin constant region. A possible multimerization domain useful in the present invention is disclosed in PCT/US90/06849 in which hybrid immunoglobulins are described. In addition a multimerization region may be engineered such that steric interactions not only promote stable interaction, but further promote the formation of heterodimers over homodimers from a mixture of monomers. See, for example, PCT/US96/01598 in which a "protuberance-into-cavity" strategy is disclosed for an interface between a first and second polypeptide for hetero-oligomerization. "Protuberances" are constructed by replacing small amino acid side chains from the interface of the first polypeptide with larger side chains (e.g. tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the protuberances are optionally created on the interface of the second polypeptide by replacing large amino acid side chains with smaller ones (e.g. alanine or threonine). The immunoglobulin sequence preferably, but not necessarily, is an immunoglobulin constant domain. The immunoglobulin moiety in the chimeras of the present invention may be obtained from $\lg G_1$, $\lg G_2$, $\lg G_3$ or $\lg G_4$, $\lg G_5$, $\lg G_6$, $\lg G_7$,

5

10

20

25

30

40

45

50

55

[0036] By "free thiol-containing compound" is meant a compound that can be incorporated into or reacted with an amino acid of a polypeptide interface of the invention such that the free thiol moiety of the compound is positioned to interact with a free thiol of moiety at the interface of additional polypeptide of the invention to form a disulfide bond. Preferably, the free thiol-containing compound is cysteine.

[0037] The term "epitope tagged" when used herein refers to a chimeric polypeptide comprising the entire chimeric heteroadhesin, or a fragment thereof, fused to a "tag polypeptide". The tag polypeptide has enough residues to provide an epitope against which an antibody can be made, yet is short enough such that it does not interfere with activity of the chimeric heteroadhesin. The tag polypeptide preferably is fairly unique so that the antibody thereagainst does not substantially cross-react with other epitopes. Suitable tag polypeptides generally have at least 6 amino acid residues and usually between about 8-50 amino acid residues (preferably between about 9-30 residues). An embodimentof the invention encompasses achimeric heteroadhesin linked to an epitope tag, which tag is used to detect the adhesin in a sample or recover the adhesin from a sample.

[0038] As used herein, "common light chain" or "common amino acid sequence of the light chain" refers to the amino acid sequence of the light chain in the multispecific antibody of the invention. Panels of antibodies were generated against at least two different antigens by panning a phage display library such as that described by Vaughan, *et al.* (1996) *supra*. The light chain sequences were compared with respect to the variable light chain amino acid sequences. Useful light chains from the compared panels are those having amino acid sequence identity of at least 80%, preferably at least 90%, more preferably at least 95%, and most preferably 100% identity. A common light chain sequence is a sequence designed to be an approximation of the two compared light chain sequences. Where the compared light chains are 100% sequence identical at the amino acid level, the common light chain is identical to the light chains from the selected library clones, even though the light chain functions in a different binding domain of the multispecific antibody. Where the compared light chains differ as described above, the common light chain may differ from one or the other, or both, of the compared light chains from the library clones. In a case in which the common light chain differs from one or the other, or both of the library clones, it is preferred that the differing residues occur outside of the antigen binding CDR residues may be determined according to a sequence definition (Kabat *et al.* (1991) *supra*) or structural definition (Chothia and Lesk (1987) J. Mol. Biol. 196:901-917).

[0039] As used herein, "amino acid sequence identity" refers to the percentage of the amino acids of one sequence are the same as the amino acids of a second amino acid sequence. 100% sequence identity between polypeptide chains means that the chains are identical.

[0040] As used herein, "polypeptide" refers generally to peptides and proteins having more than about ten amino acids. Preferably, mammalian polypeptides (polypeptides that were originally derived from a mammalian organism) are used, more preferably those which are directly secreted into the medium. Examples of bacterial polypeptides include, e.g., alkaline phosphatase and β-lactamase. Examples of mammalian polypeptides include molecules such as renin, a growth hormone, including human growth hormone; bovine growth hormone; growth hormone releasing factor; parathyroidhormone; thyroid stimulating hormone; lipoproteins; alpha-1-antitrypsin; insulin A-chain; insulin B-chain; proinsulin; follicle stimulating hormone; calcitonin; luteinizing hormone; glucagon; clotting factors such as factor VIIIC, factor IX, tissue factor, and von Willebrands factor; anti-clotting factors such as Protein C; atrial natriuretic factor; lung surfactant; a plasminogen activator, such as urokinase or human urine or tissue-type plasminogen activator (t-PA); bombesin; thrombin; hemopoietic growth factor; tumor necrosis factor-alpha and -beta; enkephalinase; RANTES (regulated on activation normally T-cell expressed and secreted); human macrophage inflammatory protein (MIP-1-alpha); a serum albumin such as human serum albumin; Muellerian-inhibiting substance; relaxin A-chain; relaxin B-chain; prorelaxin; mouse gonadotropin-associatedpeptide; a microbial protein, such as beta-lactamase; DNase; inhibin; ac-

tivin; vascularendothelialgrowth factor(VEGF); receptors for hormones or growth factors; integrin; protein A or D; rheumatoid factors; a neurotrophic factor such as bone-derived neurotrophic factor (BDNF), neurotrophin-3, -4, -5, or -6 (NT-3, NT-4, NT-5, or NT-6), or a nerve growth factor such as NGF-β; platelet-derived growth factor (PDGF); fibroblast growth factor such as aFGF and bFGF; epidermal growth factor (EGF); transforming growth factor (TGF) such as TGF-alpha and TGF-beta, including TGF-β1, TGF-p2, TGF-β3, TGF-β4, or TGF-β5; insulin-like growth factor-I and -II (IGF-I and IGF-II); des(1-3)-IGF-I(brain IGF-I), insulin-like growth factor binding proteins; CD proteins such as CD-3, CD-4, CD-8, and CD-19; erythropoietin; osteoinductive factors; immunotoxins;a bone morphogenetic protein (BMP); an interferon such as interferon-alpha,-beta, and -gamma; colony stimulating factors (CSFs), *e.g.*, M-CSF, GM-CSF, and G-CSF; interleukins (ILs), *e.g.*, IL-1 to IL-10; superoxide dismutase; T-cell receptors; surface membrane proteins; decay accelerating factor; viral antigen such as, for example, a portion of the AIDS envelope; transport proteins; homing receptors; addressins; regulatory proteins; antibodies; and fragments of any of the above-listed polypeptides.

5

10

20

25

30

45

50

[0041] The "first polypeptide" is any polypeptide which is to be associated with a second polypeptide. The first and second polypeptide meet at an "interface" (defined below). In addition to the interface, the first polypeptide may comprise one or more additional domains, such as "binding domains" (e.g. an antibody variable domain, receptor binding domain, ligand binding domain or enzymatic domain) or antibody constant domains (or parts thereof) including C_H2, C_H1 and C_L domains. Normally, the first polypeptidewill comprise at least one domain which is derived from an antibody. This domain conveniently is a constant domain, such as the C_H3 domain of an antibody and can form the interface of the first polypeptide. Exemplary first polypeptides include antibody heavy chain polypeptides, chimeras combining an antibody constant domain with a binding domain of a heterologouspolypeptide (i.e. an immunoadhesin, see definition below), receptor polypeptides (especially those which form dimers with another receptor polypeptide, e.g., interleukin-8 receptor (IL-8R) and integrin heterodimers (e.g. LFA-1 or GPIIIb/IIIa)), ligand polypeptides (e.g. nerve growth factor (NGF), neurotrophin-3 (NT-3), and brain-derived neurotrophic factor (BDNF) - see Arakawa et al. J. Biol. Chem. 269 (45): 27833-27839(1994) and Radziejewski et al. Biochem. 32(48): 1350 (1993)) and antibody variable domain polypeptides (e.g. diabodies). The preferred first polypeptide is selected from an antibody heavy chain fused to a constantdomain of an immunoglobulin, wherein the constantdomain has been altered at the interface to promote preferential interaction with a second polypeptide of the invention.

[0042] The "second polypeptide" is any polypeptide which is to be associated with the first polypeptide via an "interface". In addition to the interface, the second polypeptide may comprise additional domains such as a "binding domain" (e.g. an antibody variable domain, receptor binding domain, ligand binding domain or enzymatic domain), or antibody constant domains (or parts thereof) including C_{H2} , C_{H1} and C_{L} domains. Normally, the second polypeptide will comprise at least one domain which is derived from an antibody. This domain conveniently is a constant region, such as the C_{H3} domain of an antibody and can form the interface of the second polypeptide. Exemplary second polypeptides include antibody heavy chain polypeptides, chimeras combining an antibody constant domain with a binding domain of a heterologous polypeptide (i.e. an immunoadhesin, see definition below), receptor polypeptides (especially those which form dimers with another receptor polypeptide, e.g., interieukin-8 receptor (IL-8R) and integrin heterodimers (e.g. LFA-1 or GPIIIb/IIIa)), ligand polypeptides (e.g. nerve growth factor (NGF), neurotrophin-3 (NT-3), and brain-derived neurotrophic factor (BDNF) - see Arakawa et al. J. Biol. Chem. 269(45):27833-27839 (1994) and Radziejewski et al. Biochem. 32(48):1350(1993))and antibody variabledomain polypeptides (e.g. diabodies). The preferred second polypeptide is selected from an antibody heavy chain fused to a constant domain of an immunoglobulin, wherein the constant domain has been altered at the interface to promote preferential interaction with a first polypeptide of the invention.

[0043] A "binding domain" comprises any region of a polypeptide which is responsible for selectively binding to a molecule interest (e.g. an antigen, ligand, receptor, substrate or inhibitor). Exemplary binding domains include an antibody variable domain, receptor binding domain, ligand binding domain and an enzymatic domain. In preferred embodiments, the binding domain includes an immunoglobulin heavy chain and light chain. According to the bispecific antibodies of the invention and the method of making them, the light chain for each binding domain of the bispecific antibody is a common light chain, thereby avoiding the formation of undesired hetermultimers in which mispairing of heavy and light chains occurs.

[0044] The term "antibody" as it refers to the invention shall mean a polypeptide containing one or more domains that bind an epitope on an antigen of interest, where such domain(s) are derived from or have sequence identity with the variable region of an antibody. Examples of antibodies include full length antibodies, antibody fragments, single chain molecules, bispecific or bifunctional molecules, diabodies, chimeric antibodies (e.g. humanized and PRIMA-TIZED™ antibodies), and immunoadhesins. "Antibody fragments" include Fv, Fv', Fab, Fab', and F(ab')₂ fragments. [0045] "Humanized" forms of non-human (e.g. rodent or primate) antibodies are specific chimeric immunoglobulins, immunoglobul lin chains or fragments thereof which contain minimal sequencederived from non-human immunoglobulin. For the most part, humanized antibodies are human immunoglobulins(recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat, rabbit or primate having the desired specificity, affinity and capacity. In some instances, Fv framework region (FR) residues of the human immunoglobulin are replaced by corresponding non-

human residues. Furthermore, the humanized antibody may comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. These modifications are made to further refine and maximize antibody performance. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin sequence. The humanized antibody preferably also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin. The humanized antibody includes a PRIMATIZED™ antibody wherein the antigenbinding region of the antibody is derived from an antibody produced by immunizing macaque monkeys with the antigen of interest.

5

10

15

20

25

30

40

45

50

[0046] A "multispecificantibody" is a molecule having binding specificities for at least two different antigens. While such molecules normally will only bind two antigens (i.e. bispecific antibodies, BsAbs), antibodies with additional specificities such as trispecific antibodies are encompassed by this expression when used herein. Examples of BsAbs include those with one arm directed against a tumor cell antigen and the other arm directed against a cytotoxic trigger molecule such as anti-FcγRI/anti-CD15, anti-p185HER2/FcγRIII (CD16), anti-CD3/anti-malignant B-cell (1D10), anti-CD3/anti-p185HER2, anti-CD3/anti-p97, anti-CD3/anti-renal cell carcinoma, anti-CD3/anti-OVCAR-3, anti-CD3/L-D1 (anti-colon carcinoma), anti-CD3/anti-melanocyte stimulating hormone analog, anti-EGF receptor/anti-CD3, anti-CD3/anti-CAMA1, anti-CD3/anti-CD19, anti-CD3/MoV18, anti-neural cell ahesion molecule (NCAM)/anti-CD3, antifolate binding protein (FBP)/anti-CD3, anti-pan carcinoma associated antigen (AMOC-31)/anti-CD3; BsAbs with one arm which binds specifically to a tumor antigen and one arm which binds to a toxin such as anti-saporin/anti-ld-1, anti-CD22/anti-saporin, anti-CD7/anti-saporin, anti-CD38/anti-saporin, anti-CEA/anti-ricin A chain, anti-interferon-α (IFN-α)/anti-hybridoma idiotype, anti-CEA/anti-vinca alkaloid; BsAbs for converting enzyme activated prodrugs such as anti-CD30/anti-alkaline phosphatase (which catalyzes conversion of mitomycin phosphate prodrug to mitomycin alcohol); BsAbs which can be used as fibrinolytic agents such as anti-fibrin/anti-tissue plasminogen activator (tPA), anti-fibrin/anti-urokinase-type plasminogen activator (uPA); BsAbs for targeting immune complexes to cell surface receptors such as anti-low density lipoprotein (LDL)/anti-Fcreceptor (e.g. FcγRI, FcγRII or FcγRIII); BsAbs for use in therapy of infectious diseases such as anti-CD3/anti-herpes simplex virus (HSV), anti-T-cell receptor:CD3 complex/ anti-influenza, anti-FcγR/anti-HIV; BsAbs for tumor detection in vitro or in vivo such as anti-CEA/anti-EOTUBE, anti-CEA/anti-DPTA, anti-p185HER2/anti-hapten; BsAbs as vaccine adjuvants (see Fanger et al., supra); and BsAbs as diagnostic tools such as anti-rabbit IgG/anti-ferritin, anti-horse radish peroxidase (HRP)/anti-hormone, anti-somatostatin/anti-substance P, anti-HRP/anti-FITC, anti-CEA/anti-β-galactosidase (see Nolan et al., supra). Examples of trispecific antibodies include anti-CD3/anti-CD4/anti-CD37, anti-CD3/anti-CD5/anti-CD37 and anti-CD3/anti-CD8/anti-CD37.

[0047] As used herein, the term "immunoadhesin" designates antibody-like molecules which combine the "binding domain" of a heterologous protein (an "adhesin", *e.g.* a receptor, ligand or enzyme) with the effector functions of immunoglobulin constant domains. Structurally, the immunoadhesins comprise a fusion of the adhesin amino acid sequence with the desired binding specificity which is other than the antigen recognition and binding site (antigen combining site) of an antibody (*i.e.* is "heterologous") and an immunoglobulin constant domain sequence. The immunoglobulin constant domain sequence in the immunoadhesin may be obtained from any immunoglobulin, such as IgG_1 , IgG_2 , IgG_3 , or IgG_4 subtypes, IgA, IgE, IgD or IgM.

[0048] The term "ligand binding domain" as used herein refers to any native cell-surface receptor or any region or derivative thereof retaining at least a qualitative ligand binding ability, and preferably the biological activity of a correspondingnative receptor. In a specific embodiment, the receptor is from a cell-surface polypeptide having an extracellular domain which is homologous to a member of the immunoglobulin supergenefamily. Other typical receptors, are not members of the immunoglobulin supergenefamily but are nonetheless specifically covered by this definition, are receptors for cytokines, and in particular receptors with tyrosine kinase activity (receptor tyrosine kinases), members of the hematopoietin and nerve growth factor receptor superfamilies, and cell adhesion molecules, e. g. (E-, L- and P-) selectins.

[0049] The term "receptorbinding domain" is used to designate any native ligand for a receptor, including cell adhesion molecules, or any region or derivative of such native ligand retaining at least a qualitative receptor binding ability, and preferably the biological activity of a corresponding native ligand. This definition, among others, specifically includes binding sequences from ligands for the above-mentioned receptors.

[0050] As used herein the phrase "multispecific immunoadhesin" designates immunoadhesins (as hereinabove defined) having at least two binding specificities (*i.e.* combining two or more adhesin binding domains). Multispecific immunoadhesinscan be assembled as heterodimers, heterotrimersor heterotetramers, essentially as disclosed in WO 89/02922 (published 6 April 1989), in EP 314,317 (published 3 May 1989), and in U.S. Patent No. 5,116,964 issued 2 May 1992. Preferred muitispecific immunoadhesins are bispecific. Examples of bispecific immunoadhesins include CD4-lgG/TNFreceptor-lgG and CD4-lgG/L-selectin-lgG. The last mentioned molecule combines the lymph node binding function of the lymphocyte homing receptor (LHR, L-selectin), and the HIV binding function of CD4, and fmds

potential application in the prevention or treatment of HIV infection, related conditions, or as a diagnostic.

5

10

20

25

30

40

45

50

55

[0051] An "antibody-immunoadhesinchimera (Ab/Ia chimera)" comprises a molecule which combines at least one binding domain of an antibody (as herein defined) with at least one immunoadhesin (as defined in this application). Exemplary Ab/Ia chimeras are the bispecific CD4-IgG chimeras described by Berg *et al.*, *supra* and Chamow *et al.*, *supra*.

[0052] The "interface" comprises those "contact" amino acid residues (or other non-amino acid groups such as carbohydrate groups, NADH, biotin, FAD or haem group) in the first polypeptide which interact with one or more "contact" amino acid residues (or other non-amino acid groups) in the interface of the second polypeptide. The preferred interface is a domain of an immunoglobulinsuch as a variable domain or constant domain (or regions thereof), however the interface between the polypeptides forming a heteromultimeric receptor or the interface between two or more ligands such as NGF, NT-3 and BDNF are included within the scope of this term. The preferred interface comprises the C_H3 domain of an immunoglobulin which preferably is derived from an IgG antibody and most preferably a human IgG₁ antibody.

[0053] An "original" amino acid residue is one which is replaced by an "import" residue which can have a smaller or larger side chain volume than the original residue. The import amino acid residue can be a naturally occurring or non-naturally occurring amino acid residue, but preferably is the former. "Naturally occurring" amino acid residues are those residues encoded by the genetic code and listed in Table 1 of PCT/US96/0 1598. By "non-naturally occurring" amino acid residue is meant a residue which is not encoded by the genetic code, but which is able to covalently bind adjacent amino acid residue(s) in the polypeptide chain. Examples of non-naturally occurring amino acid residues are norleucine, ornithine, norvaline, homoserine and other amino acid residue analogues such as those described in Ellman *et al.*, Meth. Enzym. 202:301-336 (1991), for example. To generate such non-naturally occurring amino acid residues, the procedures of Noren *et al.* Science 244: 182 (1989) and Ellman *et al.*, *supra* can be used. Briefly, this involves chemically activating a suppressor tRNA with a non-naturally occurring amino acid residue followed by *in vitro* transcription and translation of the RNA. The method of the instant invention involves replacing at least one original amino acid residue, but more than one original residue can be replaced. Normally, no more than the total residues in the interface of the first or second polypeptide will comprise original amino acid residues which are replaced. The preferred original residues for replacement are "buried". By "buried" is meant that the residue is essentially inaccessible to solvent. The preferred import residue is not cysteine to prevent possible oxidation or mispairing of disulfide bonds.

[0054] By "original nucleic acid" is meant the nucleic acid encoding a polypeptide of interest which can be altered to encode within the multimerization domain amino acids whose side chains interact at the interface between the first and second polypeptide promoting stable interaction between the polypeptides. Such alterationsmay generate without limitation such stable interactions as protuberance-into-cavity, non-naturally occurring disulfide bonds, leucine zipper, hydrophobic interactions, and hydrophilic interations. Preferably, the alteration is chosen which promotes specific interaction between a first and second polypeptide of interest and effectively excludes interactions that result in undesired heteromer pairing or the formation of homomers. The original or starting nucleic acid may be a naturally occurring nucleic acid or may comprise a nucleic acid which has been subjected to prior alteration (e.g. a humanized antibody fragment). By "altering" the nucleic acid is meant that the original nucleic acid is genetically engineered or mutated by inserting, deleting or replacing at least one codon encoding an amino acid residue of interest. Normally, a codon encoding an original residue is replaced by a codon encoding an import residue. Techniques for genetically modifying a DNA in this manner have been reviewed in Mutagenesis: a Practical Approach, M.J. McPherson, Ed., (IRL Press, Oxford, UK. (1991), and include site-directedmutagenesis, cassette mutagenesis and polymerase chain reaction (PCR) mutagenesis, for example.

[0055] The protuberance, cavity, or free thiol (such as a cysteine residue for disulfide bond formation) can be "introduced" into the interface of the first or second polypeptide by synthetic means, *e.g.* by recombinant techniques, *in vitro* peptide synthesis, those techniques for introducing non-naturally occurring amino acid residues previously described, by enzymatic or chemical coupling of peptides or some combination of these techniques. According, the protuberance, cavity or free thiol which is "introduced" is "non-naturally occurring" or "non-native", which means that it does not exist in nature or in the original polypeptide (*e.g.* a humanized monoclonal antibody).

[0056] Preferably the import amino acid residue for forming the protuberance has a relatively small number of "rotamers" (e.g. about 3-6). A "rotamer" is an energetically favorable conformation of an amino acid side chain. The number of rotamers of the various amino acid residues are reviewed in Ponders and Richards, J. Mol. Biol. 193:775-791 (1987).

[0057] "Isolated" heteromultimer means heteromultimer which has been identified and separated and/or recovered from a component of its natural cell culture environment. Contaminant components of its natural environment are materials which would interfere with diagnostic or therapeutic uses for the heteromultimer, and may include enzymes, hormones, and other proteinaceous or nonproteinaceous solutes. In preferred embodiments, the heteromultimer will be purified (1) to greater than 95% by weight of protein as determined by the Lowry method, and most preferably more than 99% by weight, (2) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence

by use of a spinning cup sequenator, or (3) to homogeneity by SDS-PAGE under reducing or nonreducing conditions using Coomassie blue or, preferably, silver stain.

[0058] The heteromultimers of the present invention are generally purified to substantial homogeneity. The phrases "substantially homogeneous", "substantially homogeneous form" and "substantial homogeneity" are used to indicate that the product is substantially devoid of by-products originated from undesired polypeptide combinations (*e.g.* homomultimers). Expressed in terms of purity, substantial homogeneity means that the amount of by-products does not exceed 10%, and preferably is below 5%, more preferably below 1%, most preferably below 0.5%, wherein the percentages are by weight.

[0059] The expression "control sequences" refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, a ribosome binding site, and possibly, other as yet poorly understood sequences. Eukaryotic cells are known to utilize promoters, polyadenylationsignals, and enhancers. [0060] Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, "operably linked" means that the DNA sequences being linked are contiguous and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accord with conventional practice.

II. Preparation of the Heteromultimer

5

10

20

25

30

40

45

50

55

1. Preparation of the Starting Materials

[0061] As a first step, the first and second polypeptide (and any additional polypeptides forming the heteromultimer) are selected. Normally, the nucleic acid encoding these polypeptides needs to be isolated so that it can be altered to encode the protuberance or cavity, or both, as herein defined. However, the mutations can be introduced using synthetic means, e.g. by using a peptide synthesizer. Also, in the case where the import residue is a non-naturally occurring residue, the method of Noren et al., supra is available for making polypeptides having such substitutions. Additionally, part of the heteromultimer is suitably made recombinantly in cell culture and other part(s) of the molecule are made by those techniques mentioned above.

[0062] Techniques for isolating antibodies and preparing immunoadhesins follow. However, it will be appreciated that the heteromultimer can be formed from, or incorporate, other polypeptides using techniques which are known in the art. For example, nucleic acid encoding a polypeptide of interest (e.g. a ligand, receptor or enzyme) can be isolated from a cDNA library prepared from tissue believed to possess the polypeptide mRNA and to express it at a detectable level. Libraries are screened with probes (such as antibodies or oligonucleotides of about 20-80 bases) designed to identify the gene of interest or the protein encoded by it. Screening the cDNA or genomic library with the selected probe may be conducted using standard procedures as described in chapters 10-12 of Sambrook *et al.*, Molecular Cloning: A Laboratory Manual (New York: Cold Spring Harbor Laboratory Press, 1989).

(I) Antibody preparation

[0063] Several techniques for the production of antibodies have been described which include the traditional hybridoma method for making monoclonal antibodies, recombinant techniques for making antibodies (including chimeric antibodies, *e.g.* humanized antibodies), antibody production in transgenic animals and the recently described phage display technology for preparing "fully human" antibodies. These techniques shall be described briefly below.

[0064] Polyclonal antibodies to the antigen of interest generally can be raised in animals by multiple subcutaneous (sc) or intraperitoneal (ip) injections of the antigen and an adjuvant. It may be useful to conjugate the antigen (or a fragment containing the target amino acid sequence) to a protein that is immunogenic in the species to be immunized, e.g., keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, or soybean trypsin inhibitor using a bifunctional or derivatizing agent, for example maleimidobenzoyl sulfosuccinimide ester (conjugation through cysteine residues), N-hydroxysuccinimide (through lysine residues), glutaraldehyde, succinic anhydride, SOCl₂, or R'N=C=NR, where R and R¹ are different alkyl groups. Animals are immunized against the immunogenic conjugates or derivatives by combining I mg of 1 µg of conjugate (for rabbits or mice, respectively) with 3 volumes of Freud's complete adjuvant and injecting the solution intradermally at multiple sites. One month later the animals are boosted with 1/5 to 1/10 the original amount of conjugate in Freud's complete adjuvant by subcutaneous injection at multiple sites. 7 to 14 days later the animals are bled and the serum is assayed for antibody titer. Animals are boosted until the titer plateaus.

Preferably, the animal is boosted with the conjugate of the same antigen, but conjugated to a different protein and/or through a different cross-linking reagent. Conjugates also can be made in recombinantcell culture as protein fusions. Also, aggregating agents such as alum are used to enhance the immune response.

5

10

15

20

25

30

40

45

50

55

[0065] Monoclonal antibodies are obtained from a population of substantially homogeneous antibodies using the hybridoma method first described by Kohler and Milstein, Nature 256:495 (1975) or may be made by recombinant DNA methods (Cabilly et al., U.S. Patent No. 4,816,567). In the hybridoma method, a mouse or other appropriate host animal, such as hamster, is immunized as hereinabove described to elicit lymphocytes that produce, or are capable of producing, antibodies that will specifically bind to the protein used for immunization. Alternatively, lymphocytes may be immunized in vitro. Lymphocytes then are fused with myeloma cells using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, Monoclonal Antibodies: Principles and Practice, pp.59-103 (Academic Press, 1986)). The hybridoma cells thus prepared are seeded and grown in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, parental myeloma cells. For example, if the parental myeloma cells lack the enzyme hypoxanthineguanine phosphoribosyltransferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine (HAT medium), which substancespreventthe growth of HGPRT-deficient cells. Preferred myeloma cells are those that fuse efficiently, support stable high level expression of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. Among these, preferred myeloma cell lines are murine myeloma lines, such as those derived from MOPC-21 and MPC-11 mouse tumors available from the Salk Institute Cell Distribution Center, San Diego, California USA, and SP-2 cells available from the American Type Culture Collection, Rockville, Maryland USA. Human myeloma and mouse-human heteromyelomacell lines also have been described for the production of human monoclonal antibodies (Kozbor, J. Immunol., 133:3001 (1984); and Brodeur et al., Monoclonal Antibody Production Techniques and Applications, pp.51-63, Marcel Dekker, Inc., New York, 1987). See, also, Boerner et al., J. Immunol., 147 (1):86-95 (1991) and WO 91/17769, published Nov 28, 1991, for techniques for the production of human monoclonal antibodies. Culture medium in which hybridoma cells are growing is assayed for production of monoclonal antibodies directed against the antigen of interest. Preferably, the binding specificity of monoclonal antibodies produced by hybridoma cells is determined by immunoprecipitation or by an in vitro binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA). The binding affinity of the monoclonal antibody can, for example, be determined by the Scatchard analysis of Munson and Pollard, Anal. Biochem. 107:220 (1980). After hybridoma cells are identified that produce antibodies of the desired specificity, affinity, and/or activity, the clones may be subcloned by limiting dilution procedures and grown by standard methods. Goding, Monoclonal Antibodies: Principles and Practice, pp.59-104 (Academic Press, 1986). Suitable culture media for this purpose include, for example, Dulbecco's Modified Eagle's Medium or RPM1-1640 medium. In addition, the hybridoma cells may be grown in vivo as ascites tumors in an animal. The monoclonal antibodies secreted by the subclones are suitably separated from the culture medium, ascites fluid, or serum by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

[0066] Alternatively, it is now possible to produce transgenic animals (e.g. mice) that are capable, upon immunization, of producing a full repertoire of human antibodies in the absence of endogenous immunoglobulin production. For example, it has been described that the homozygous deletion of the antibody heavy chain joining region (J_H) gene in chimeric and germ-line mutant mice results in complete inhibition of endogenous antibody production. Transfer of the human germ-line immunoglobulin gene array in such germ-line mutant mice will result in the production of human antibodies upon antigen challenge. See, *e.g.*, Jakobovits *et al.*, Proc. Natl. Acad. Sci. USA <u>90</u>:2551-255 (1993); Jakobovits *et al.*, Nature <u>362</u>:255-258 (1993); Fishwild, D.M., *et al.* (1996) Nat. Biotech <u>14</u>:845-851; and Mendez. M. J., *et al.* (1997) Nat. Genetics 15:146-156).

[0067] In a further embodiment, antibodies or antibody fragments can be isolated from antibody phage libraries generated using the techniques described in McCafferty *et al.*, Nature, 348:552-554 (1990), using the antigen of interest to select for a suitable antibody or antibody fragment. Clackson *et al.*, Nature, 352:624-628 (1991) and Marks *et al.*, J. Mol. Biol., 222:581-597 (1991) describe the isolation of murine and human antibodies, respectively, using phage libraries. Subsequent publications describe the production of high affinity (nM range) human antibodies by chain shuffling (Mark *et al.*, Bio/Technol. 10:779-783 (1992)), as well as combinatorial infection and *in vivo* recombination as a strategy for constructing very large phage libraries (Waterhouse *et al.*, Nuc. Acids Res., 21:2265-2266 (1993); Griffiths, A. D., *et al.* (1994) EMBO J. 13:3245-3260; and Vaughan, *et al.* (1996) *supra*). Thus, these techniques are viable alternatives to traditional monoclonal antibody hybridoma techniques for isolation of "monoclonal" antibodies (especially human antibodies) which are encompassed by the present invention.

[0068] DNA encoding the antibodies of the invention is readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein,

to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also may be modified, for example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences, Morrison *et al.*, Proc. Nat. Acad. Sci. <u>81</u>:6851 (1984). In that manner, "chimeric" antibodies are prepared that have the binding specificity of an anti-antigen monoclonal antibody herein.

[0069] Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. Humanization can be performed essentially following the method of Winter and co-workers (Jones et al., Nature 321:522-525 (1986); Riechmann et al., Nature 332:323-327 (1988); Verhoeyen et al., Science 239:1534-1536 (1988)), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such "humanized" antibodies are chimeric antibodies (Cabilly, supra), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues, and possibly some FR residues, are substituted by residues from analogous sites in rodent antibodies. It is important that antibodies be humanized with retention of high affinity for the antigen and other favorable biological properties. To achieve this goal, according to a preferred method, humanized antibodies are prepared by a process of analysis of the parental sequences and various conceptual humanized products using three dimensional models of the parental and humanized sequences. Three dimensional immunoglobulin models are familiar to those skilled in the art. Computer programs are available which illustrate and display probable threedimensional conformational structures of selected candidate immunoglobulin sequences. Inspection of these displays permits analysis of the likely role of the residues in the functioning of the candidate immunoglobulin sequence, i.e., the analysis of residues that influence the ability of the candidate immunoglobulin to bind its antigen. In this way, FR residues can be selected and combined from the consensus and import sequence so that the desired antibody characteristic, such as increased affinity for the target antigen(s), is achieved. For further details see WO 92/22653, published Dec 23, 1992.

25 (ii) Immunoadhesin preparation

5

10

15

20

30

45

50

[0070] Immunoglobulins (Ig) and certain variants thereof are known and many have been prepared in recombinant cell culture. For example, see U.S. Patent No. 4,745,055; EP 256,654; Faulkner *et al.*, Nature 298:286 (1982); EP 120,694; EP 125,023; Morrison, J. Immun. 123:793 (1979); Köhler *et al.*, Proc. Natl. Acad. Sci. USA 77:2197 (1980); Raso *et al.*, Cancer Res. 41:2073 (1981); Morrison *et al.*, Ann. Rev. Immunol. 2:239 (1984); Morrison, Science 229: 1202 (1985); Morrison *et al.*, Proc. Natl. Acad. Sci. USA 81:6851 (1984); EP 255,694; EP 266,663; and WO 88/03559. Reassorted immunoglobulin chains also are known. See, for example, U.S. Patent No. 4,444,878; WO 88/03565; and EP 68,763 and references cited therein.

[0071] Chimeras constructed from an adhesin binding domain sequence linked to an appropriate immunoglobulin constant domain sequence (immunoadhesins) are known in the art. Immunoadhesinsreported in the literature include fusions of the T cell receptor (Gascoigne *et al.*, Proc. Natl. Acad. Sci. USA 84:2936-2940 (1987)); CD4 (Capon *et al.*, Nature 337:525-531 (1989); Traunecker *et al.*, Nature 339:68-70 (1989); Zettmeissl *et al.*, DNA Cell Biol. USA 9:347-353 (1990); and Byrn *et al.*, Nature 344:667-670 (1990)); L-selectin (homing receptor) (Watson *et al.*, J. Cell. Biol. 110: 2221-2229 (1990); and Watson *et al.*, Nature 349:164-167 (1991)); CD44 (Aruffo *et al.*, Cell 61:1303-1313 (1990)); CD28 and B7 (Linsley *et al.*, J. Exp. Med. 173:721-730 (1991)); CTLA-4 (Lisley *et al.*, J. Exp. Med. 174:561-569 (1991)); CD22 (Stamenkovic *et al.*, Cell 66:1133-1144 (1991)); TNF receptor (Ashkenazi *et al.*, Proc. Natl. Acad. Sci. USA 88: 10535-10539 (1991); Lesslauer *et al.*, Eur. J. Immunol. 27:2883-2886 (1991); and Peppel *et al.*, J. Exp. Med. 174: 1483-1489(1991)); and IgE receptor α (Ridgwayand Gorman, J. Cell. Biol. Vol. 115, Abstract No. 1448 (1991)).

[0072] The simplest and most straightforward immunoadhesin design combines the binding domain(s) of the adhesin (e.g. the extracellular domain (ECD) of a receptor) with the hinge and Fc regions of an immunoglobu lin heavy chain. Ordinarily, when preparing the immunoadhesins of the present invention, nucleic acid encoding the binding domain of the adhesin will be fused C-terminally to nucleic acid encoding the N-terminus of an immunoglobulin constant domain sequence, however N-terminal fusions are also possible.

[0073] Typically, in such fusions the encoded chimeric polypeptide will retain at least functionally active hinge, $C_{H}2$ and $C_{H}3$ domains of the constant region of an immunoglobulin heavy chain. Fusions are also made to the C-terminus of the Fc portion of a constant domain, or immediately N-terminal to the $C_{H}1$ of the heavy chain or the corresponding of the light chain. The precise site at which the fusion is made is not critical; particular sites are well known and may be selected in order to optimize the biological activity, secretion, or binding characteristics of the 1a.

[0074] In a preferred embodiment, the adhesin sequence is fused to the N-terminus of the Fc domain of immunoglobu lin G₁ (IgG₁). It is possible to fuse the entire heavy chain constant region to the adhesin sequence. However, more preferably, a sequence beginning in the hinge region just upstream of the papain cleavage site which defines IgG Fc chemically (*i.e.* residue 216, taking the first residue of heavy chain constant region to be 114), or analogous sites of other immunoglobulins is used in the fusion. In a particularly preferred embodiment, the adhesin amino acid sequence

is fused to (a) the hinge region and $C_{H}2$ and $C_{H}3$ or (b) the $C_{H}1$, hinge, $C_{H}2$ and $C_{H}3$ domains, of an IgG_{1} , IgG_{2} , or IgG_{3} heavy chain. The precise site at which the fusion is made is not critical, and the optimal site can be determined by routine experimentation.

[0075] For bispecific immunoadhesins, the immunoadhesins are assembled as multimers, and particularly as heterodimers or heterotetramers. Generally, these assembled immunoglobulins will have known unit structures. A basic four chain structural unit is the form in which IgG, IgD, and IgE exist. A four chain unit is repeated in the higher molecular weight immunoglobulins;IgM generally exists as a pentamer of four basic units held together by disulfide bonds. IgA globulin, and occasionally IgG globulin, may also exist in multimeric form in serum. In the case of multimer, each of the four units may be the same or different.

[0076] Various exemplary assembled immunoadhesins within the scope herein are schematically diagrammed below:

```
 \begin{array}{l} \text{(a) } \mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{L}}; \\ \text{(b) } \mathsf{AC}_{\mathsf{H}}\text{-}[\mathsf{AC}_{\mathsf{H}},\,\mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}},\,\mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{V}_{\mathsf{H}}\mathsf{C}_{\mathsf{H}},\,\mathsf{or}\,\,\mathsf{V}_{\mathsf{L}}\mathsf{C}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}}]; \\ \text{(c) } \mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}}\text{-}[\mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}},\,\mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{V}_{\mathsf{H}}\mathsf{C}_{\mathsf{H}},\,\mathsf{V}_{\mathsf{L}}\mathsf{C}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}},\,\mathsf{or}\,\,\mathsf{V}_{\mathsf{L}}\mathsf{C}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}}]; \\ \text{(d) } \mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{V}_{\mathsf{H}}\mathsf{C}_{\mathsf{H}},\,\mathsf{or}\,\,\mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{V}_{\mathsf{H}}\mathsf{C}_{\mathsf{H}},\,\mathsf{or}\,\,\mathsf{V}_{\mathsf{L}}\mathsf{C}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}}]; \\ \text{(e) } \mathsf{V}_{\mathsf{L}}\mathsf{C}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}}\text{-}[\mathsf{AC}_{\mathsf{L}}\text{-}\mathsf{V}_{\mathsf{H}}\mathsf{C}_{\mathsf{H}},\,\mathsf{or}\,\,\mathsf{V}_{\mathsf{L}}\mathsf{C}_{\mathsf{L}}\text{-}\mathsf{AC}_{\mathsf{H}}];\,\mathsf{and} \\ \text{(f) } [\mathsf{A}\text{-}\mathsf{Y}]_{\mathsf{n}}\text{-}[\mathsf{V}_{\mathsf{L}}\mathsf{C}_{\mathsf{L}}\text{-}\mathsf{V}_{\mathsf{H}}\mathsf{C}_{\mathsf{H}}]_{2}, \\ \end{array}
```

wherein each A represents identical or different adhesin amino acid sequences;

V_I is an immunoglobulin light chain variable domain;

V_H is an immunoglobulin heavy chain variable domain;

C_L is an immunoglobulin light chain constant domain;

C_H is an immunoglobulin heavy chain constant domain;

n is an integer greater than 1;

5

10

25

30

45

50

55

Y designates the residue of a covalent cross-linking agent.

[0077] In the interests of brevity, the foregoing structures only show key features; they do not indicate joining (J) or other domains of the immunoglobulins, nor are disulfide bonds shown. However, where such domains are required for binding activity, they shall be constructed to be present in the ordinary locations which they occupy in the immunoglobulin molecules.

[0078] Alternatively, the adhesin sequences can be inserted between immunoglobulin heavy chain and light chain sequences, such that an immunoglobulin comprising a chimeric heavy chain is obtained. In this embodiment, the adhesin sequences are fused to the 3' end of an immunoglobulin heavy chain in each arm of an immunoglobulin, either between the hinge and the C_H^2 domain, or between the C_H^2 and C_H^3 domains. Similar constructs have been reported by Hoogenboom, *et al.*, Mol. Immunol. <u>28</u>:1027-1037 (1991).

[0079] An immunoglobulin light chain might be present either covalently associated to an adhesin-immunoglobulin heavy chain fusion polypeptide, or directly fused to the adhesin. In the former case, DNA encoding an immunoglobulin light chain is typically coexpressed with the DNA encoding the adhesin-immunoglobulin heavy chain fusion protein. Upon secretion, the hybrid heavy chain and the light chain will be covalently associated to provide an immunoglobulin-like structure comprising two disulfide-linked immunoglobulin heavy chain-light chain pairs. Methods suitable for the preparation of such structures are, for example, disclosed in U.S. Patent No. 4,816,567, issued 28 March 1989.

[0080] In a preferred embodiment, the immunoglobulin sequences used in the construction of the immunoadhesins of the present invention are from an $\lg G$ immunoglobulin heavy chain constant domain. For human immunoadhesins, the use of human $\lg G_1$ and $\lg G_3$ immunoglobulin sequences is preferred. A major advantage of using $\lg G_1$ is that $\lg G_1$ immunoadhesins can be purified efficiently on immobilized protein A. In contrast, purification of $\lg G_3$ requires protein G, a significantly less versatile medium. However, other structural and functional properties of immunoglobulins should be considered when choosing the \lg fusion partner for a particular immunoadhesinconstruction. For example, the $\lg G_3$ hinge is longer and more flexible, so it can accommodate larger "adhesin" domains that may not fold or function properly when fused to $\lg G_1$. Another consideration may be valency; $\lg G$ immunoadhesins are bivalent homodimers, whereas \lg subtypes like $\lg A$ and $\lg M$ may give rise to dimeric or pentameric structures, respectively, of the basic \lg homodimer unit. For immunoadhesins designed for *in vivo* application, the pharmacokinetic properties and the effector functions specified by the Fc region are important as well. Although $\lg G_1$, $\lg G_2$ and $\lg G_4$ all have *in vivo* half-lives of 21 days, their relative potencies at activating the complement system are different. $\lg G_4$ does not activate complement, and $\lg G_2$ is significantly weaker at complement activation than $\lg G_1$. Moreover, unlike $\lg G_1$, $\lg G_2$ does not bind to Fc receptors on mononuclear cells or neutrophils. While $\lg G_3$ is optimal for complement activation, its *in vivo* half-life is approximately one third of the other $\lg G$ isotypes. Another important consideration for immunoadhesins designed to

be used as human therapeutics is the number of allotypic variants of the particular isotype. In general, IgG isotypes with fewer serologically-defined allotypic sites, two of which (G1m and 2) are located in the Fc region; and one of these sites, G1m1, is non-immunogenic. In contrast, there are 12 serologically-defined allotypes in IgG3, all of which are in the Fc region; only three of these sites (G3m5, 11 and 21) have one allotype which is nonimmunogenic. Thus, the potential immunogenicity of a γ 3 immunoadhesin is greater than that of a γ 1 immunoadhesin.

[0081] Immunoadhesinsare most conveniently constructed by fusing the cDNA sequence encoding the adhesin portion in-frame to an 1g cDNA sequence. However, fusion to genomic 1g fragments can also be used (see, e.g. Gascoigneet *al.*, supra; Aruffo *et al.*, Cell <u>61</u>:1303-1313 (1990); and Stamenkovic *et al.*, Cell <u>66</u>:1133-1144 (1991)). The latter type of fusion requires the presence of Ig regulatory sequences for expression. cDNAs encoding IgG heavy-chain constant regions can be isolated based on published sequences from cDNA libraries derived from spleen or peripheral blood lymphocytes, by hybridization or by polymerase chain reaction (PCR) techniques. The cDNAs encoding the "adhesin" and the Ig parts of the immunoadhesin are inserted in tandem into a plasmid vector that directs efficient expression in the chosen host cells.

2. Generating Protuberance and/or Cavity

[0082] As a first step to selecting original residues for forming the protuberance and/or cavity, the three-dimensional structure of the heteromultimeris obtained using techniques which are well known in the art such as X-ray crystallog-raphy or NMR. Based on the three-dimensional structure, those skilled in the art will be able to identify the interface residues.

[0083] The preferred interface is the C_H3 domain of an immunoglobulin constant domain. The interface residues of the C_H3 domains of IgG, IgA, IgD, IgE and IgM have been identified (see, for example, PCT/US96/01598), including those which are optimal for replacing with import residues; as were the interface residues of various IgG subtypes and "buried" residues. The basis for engineering the C_H3 interface is that X-ray crystallography has demonstrated that the intermolecu lar association between human IgG₁ heavy chains in the Fc region includes extensive protein/protein interaction between C_H3 domains whereas the glycosylated C_H2 domains interact via their carbohydrate (Deisenhofer, Biochem. 20:2361-2370 (1981)). In addition there are two inter-heavy chain disulfide bonds which are efficiently formed during antibody expression in mammalian cells unless the heavy chain is truncated to remove C_H2 and C_H3 domains (King et al., Biochem. J. 281:317 (1992)). Thus, heavy chain assembly appears to promote disulfide bond formation rather than vice versa. Taken together these structural and functionaldata led to the hypothesis that antibody heavy chain association is directed by the C_H3 domains. It was further speculated that the interface between C_H3 domains might be engineered to promote formation of heteromultimers of different heavy chains and hinder assembly of correspondinghomomultimers. The experiments described herein demonstrated that it was possible to promote the formation of heteromultimers over homomultimers using this approach. Thus, it is possible to generate a polypeptide fusion comprising a polypeptide of interest and the C_H3 domain of an antibody to form a first or second polypeptide. The preferred C_H3 domain is derived from an IgG antibody, such as an human IgG₁.

[0084] Those interface residues which can potentially constitute candidates for forming the protuberance or cavity are identified. It is preferable to select "buried" residues to be replaced. To determine whether a residue is buried, the surface accessibility program of Lee *et al.* J. Mol. Biol. <u>55</u>:379-400 (1971) can be used to calculate the solvent accessibility (SA) of residues in the interface. Then, the SA for the residues of each of the first and second polypeptide can be separately calculated after removal of the other polypeptide. The difference in SA of each residue between the monomer and dimer forms of the interface can then be calculated using the equation: SA (dimer) - SA (monomer). This provides a list of residues which lose SA on formation of the dimer. The SA of each residue in the dimer is compared to the theoretical SA of the same amino acid in the tripeptide Gly-X-Gly, where X = the amino acid of interest (Rose *et al.* Science <u>229</u>:834-838 (1985)). Residues which (a) lost SA in the dimer compared to the monomer and (b) had an SA less than 26% of that in their corresponding tripeptide are considered as interface residues. Two categories may be delineated: those which have an SA < 10% compared to their corresponding tripeptide (*i.e.* "buried") and those which have 25% > SA > 10% compared to their corresponding tripeptide (*i.e.* "partially buried") (see Table 1, below).

55

50

45

5

10

15

20

25

30

TABLE I

5	

	SA Lost Mond	omer → Dimer	% Trip	peptide
Residue No. [†]	Polypeptide A	Polypeptide B	Polypeptide A	Polypeptide B
Q347	22.1	31.0	25.0	26.5
Y349	79.8	83.9	5.2	5.7
L351	67.4	77.7	3.9	2.0
S354	53.4	52.8	11.3	11.7
E357	43.7	45.3	0.4	1.3
S364	21.5	15.1	0.5	1.4
Т366	29.3	25.8	0.0	0.1
L368	25.5	29.7	1.0	1.1
K370	55.8	62.3	11.5	11.0
T394	64.0	58.5	0.6	1.4
V397	50.3	49.5	13.2	11.0
D399	39.7	33.7	5.7	5.7
F405	53.7	52.1	0.0	0.0
Y407	89.1	90.3	0.0	0.0
K409	86.8	92.3	0.7	0.6
T411	4.3	7.5	12.7	9.8

[†] residue numbering as in IgG crystal structure (Deisenhofer, Biochemistry 20:2361-2370 (1981)).

[0085] The effect of replacing residues on the polypeptide chain structure can be studied using a molecular graphics modeling program such as the Insight™ program (Biosym Technologies). Using the program, those buried residues in the interface of the first polypeptide which have a small side chain volume can be changed to residues having a larger side chain volume (*i.e.* a protuberance), for example. Then, the residues in the interface of the second polypeptide which are in proximity to the protuberance are examined to find a suitable residue for forming the cavity. Normally, this residue will have a large side chain volume and is replaced with a residue having a smaller side chain volume. In certain embodiments, examination of the three-dimensional structure of the interface will reveal a suitably positioned and dimensioned protuberance on the interface of the first polypeptide or a cavity on the interface of the second polypeptide. In these instances, it is only necessary to model a single mutant, *i.e.*, with a synthetically introduced protuberance or cavity.

[0086] With respect to selecting potential original residues for replacement where the first and second polypeptide each comprise a C_H3 domain, the C_H3/C_H3 interface of human IgG_1 involves sixteen residues on each domain located on four anti-parallel β -strands which buries 1090 Å² from each surface (Deisenhofer, *supra*) and Miller, J. Mol. Biol. 216:965 (1990)). Mutations are preferably targeted to residues located on the two central anti-parallel β -strands. The aim is to minimize the risk that the protuberances which are created can be accommodated by protruding into surrounding solvent rather than by compensatory cavities in the partner C_H3 domain.

[0087] Once the preferred original/import residues are identified by molecular modeling, the amino acid replacements are introduced into the polypeptide using techniques which are well known in the art. Normally the DNA encoding the polypeptide is genetically engineered using the techniques described in Mutagenesis: a Practical Approach, *supra*.

[0088] Oligonucleotide-mediated mutagenesis is a preferred method for preparing substitution variants of the DNA encoding the first or second polypeptide. This technique is well known in the art as described by Adelman *et al.*, DNA, 2:183 (1983). Briefly, first or second polypeptide DNA is altered by hybridizing an oligonucleotideencodingthe desired mutation to a DNA template, where the template is the single-stranded form of a plasmid or bacteriophage containing the unaltered or native DNA sequence of heteromultimer. After hybridization, a DNA polymerase is used to synthesize an entire second complementary strand of the template that will thus incorporate the oligonucleotide primer, and will code for the selected alteration in the heteromultimer DNA.

[0089] Cassette mutagenesis can be performed as described Wells *et al.* Gene <u>34</u>:315 (1985) by replacing a region of the DNA of interest with a synthetic mutant fragment generated by annealing complimentary oligonucleotides. PCR mutagenesis is also suitable for making variants of the first or second polypeptide DNA. While the following discussion refers to DNA, it is understood that the technique also finds application with RNA. The PCR technique generally refers to the following procedure (see Erlich, Science, 252:1643-1650 (1991), the chapter by R. Higuchi, p. 61-70).

[0090] This invention also encompasses, in addition to the protuberance or cavity mutations, amino acid sequence variants of the heteromultimer which can be prepared by introducing appropriate nucleotide changes into the heteromultimer DNA, or by synthesis of the desired heteromultimer polypeptide. Such variants include, for example, deletions from, or insertions or substitutions of, residues within the amino acid sequences of the first and second polypeptides forming the heteromultimer. Any combination of deletion, insertion, and substitution is made to arrive at the final construct, provided that the final construct possesses the desired antigen-binding characteristics. The amino acid changes also may alter post-translational processes of the heteromultimer, such as changing the number or position of glycosylation sites.

[0091] A useful method for identification of certain residues or regions of the heteromultimer polypeptides that are preferred locations for mutagenesis is called "alanine scanning mutagenesis," as described by Cunningham and Wells, Science, 244:1081-1085 (1989). Here, a residue or group of target residues are identified (e.g. charged residues such as arg, asp, his, lys, and glu) and replaced by a neutral or negatively charged amino acid (most preferably alanine or polyalanine) to affect the interaction of the amino acids with the surrounding aqueous environment in or outside the cell. Those domains demonstrating functional sensitivity to the substitutions then are refined by introducing further or other variants at or for the sites of substitution. Thus, while the site for introducing an amino acid sequence variation is predetermined, the nature of the mutation *per se* need not be predetermined.

[0092] Normally the mutations will involve conservative amino acid replacements in non-functional regions of the heteromultimer. Exemplary mutations are shown in Table 2.

Table 2

5	Original Residue	Exemplary Substitutions	Preferred Substitutions
	Ala (A)	Val; Leu; He	Val
	Arg (R)	Lys; Gln; Asn	Lys
10	Asn (N)	Gln; His; Lys; Arg	Gln
	Asp (D)	Glu	Glu
	Cys (C)	Ser	Ser
15	Gln (Q)	Asn	Asn
	Glu (E)	Asp	Asp
20	Gly (G)	Pro; Ala	Ala
	His (H)	Asn; Gln; Lys; Arg	Arg
	Ile (I)	Leu; Val; Met; Ala; Phe; Norleucine	Leu
25	Leu (L)	Norleucine; Ile; Val; Met; Ala; Phe	Ile
	Lys (K)	Arg; Gln; Asn	Arg
30	Met (M)	Leu; Phe; Ile	Leu
	Phe (F)	Leu; Val; Ile; Ala; Tyr	Leu
25	Pro (P)	Ala	Ala
35	Ser (S)	Thr	Thr
	Thr (T)	Ser	Ser
40	Trp (W)	Tyr; Phe	Tyr
	Tyr (Y)	Trp; Phe; Thr; Ser	Phe
	Val (V)	Ile; Leu; Met; Phe; Ala;	Leu

Norleucine

45

50

55

[0093] Covalent modifications of the heteromultimer polypeptides are included within the scope of this invention. Covalent modifications of the heteromultimer can be introduced into the molecule by reacting targeted amino acid residues of the heteromultimer or fragments thereof with an organic derivatizing agent that is capable of reacting with selected side chains or the N- or C-terminal residues. Another type of covalent modification of the heteromultimer-polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. By altering is meant deleting one or more carbohydrate moieties found in the original heteromultimer, and/or adding one or more glycosylation sites that are not present in the original heteromultimer. Addition of glycosylation sites to the heteromultimer polypeptide is conveniently accomplished by altering the amino acid sequence such that it contains one or more N-linked glycosylationsites. The alteration may also be made by the addition of, or substitution by, one or more serine or threonine residues to the original heteromultimer sequence (for O-linked glycosylation sites). For ease, the heteromultimeramino acid sequence is preferably altered through changes at the DNA level, particularly

by mutating the DNA encoding the heteromultimer polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids. Another means of increasing the number of carbohydrate moieties on the heteromultimer polypeptide is by chemical or enzymatic coupling of glycosides to the polypeptide. These methods are described in WO 87/05330 published 11 September 1987, and in Aplin and Wriston, CRC Crit. Rev. Biochem., pp. 259-306 (1981). Removal of carbohydrate moieties present on the heteromultimer may be accomplished chemically or enzymatically.

[0094] Another type of covalent modification of heteromultimer comprises linking the heteromultimer polypeptide to one of a variety of nonproteinaceouspolymers, *e.g.*, polyethylene glycol, polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835;4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337.

[0095] Since it is often difficult to predict in advance the characteristics of a variant heteromultimer, it will be appreciated that some screening of the recovered variant will be needed to select the optimal variant.

3. Expression of Heteromultimer having common light chains

5

10

15

20

25

30

45

50

55

[0096] Following mutation of the DNA and selection of the common light chain as disclosed herein, the DNA encoding the molecules is expressed using recombinanttechniqueswhich are widely available in the art. Often, the expression system of choice will involve a mammalian cell expression vector and host so that the heteromultimer is appropriately glycosylated (e.g. in the case of heteromultimerscomprising antibody domains which are glycosylated). However, the molecules can also be produced in the prokaryotic expression systems elaborated below. Normally, the host cell will be transformed with DNA encoding both the first polypeptide, the second polypeptide, the common light chain polypeptide, and other polypeptide(s) required to form the heteromultimer, on a single vector or independent vectors. However, it is possible to express the first polypeptide, second polypeptide, and common light chain polypeptide (the heteromultimer components) in independent expression systems and couple the expressed polypeptides *in vitro*.

[0097] The nucleic acid(s) (e.g., cDNA or genomic DNA) encoding the heteromultimer and common light chain is inserted into a replicable vector for further cloning (amplification of the DNA) or for expression. Many vectors are available. The vector components generally include, but are not limited to, one or more of the following: a signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence.

[0098] The polypeptides of the heteromultimer components may be produced as fusion polypeptides with a signal sequence or other polypeptide having a specific cleavage site at the N-terminus of the mature protein or polypeptide. In general, the signal sequence may be a component of the vector, or it may be a part of the DNA that is inserted into the vector. The heterologous signal sequence selected preferably is one that is recognized and processed (*i.e.*, cleaved by a signal peptidase) by the host cell. For prokaryotic host cells, the signal sequence may be substituted by a prokaryotic signal sequence selected, for example, from the group of the alkaline phosphatase, penicillinase, lpp, or heat-stable enterotoxin II leaders. For yeast secretion the native signal sequence may be substituted by, *e.g.*, the yeast invertase leader, alpha factor leader (including *Saccharomyces* and *Kluyveromyces* α-factor leaders, the latter described in U.S. Pat. No. 5,010,182 issued 23 April 1991), or acid phosphatase leader, the *C. albicans* glucoamylase leader (EP 362,179 published 4 April 1990), or the signal described in WO 90/13646 published 15 November 1990. In mammalian cell expression the native signal sequence (*e.g.*, the antibody or adhesin presequence that normally directs secretion of these molecules from human cells *in vivo*) is satisfactory, although other mammalian signal sequences may be suitable as well as viral secretory leaders, for example, the herpes simplex gD signal. The DNA for such precursorregion is ligated in reading frame to DNA encoding the polypeptides forming the heteromultimer.

[0099] Both expression and cloning vectors contain a nucleic acid sequence that enables the vector to replicate in one or more selected host cells. Generally, in cloning vectors this sequence is one that enables the vector to replicate independently of the host chromosomal DNA, and includes origins of replication or autonomously replicating sequences. Such sequences are well known for a variety of bacteria, yeast, and viruses. The origin of replication from the plasmid pBR322 is suitable for most Gram-negative bacteria, the 2μ plasmid origin is suitable for yeast, and various viral origins (SV40, polyoma, adenovirus, VSV or BPV) are useful for cloning vectors in mammalian cells. Generally, the origin of replication component is not needed for mammalian expression vectors (the SV40 origin may typically be used only because it contains the early promoter).

[0100] Expression and cloning vectors should contain a selection gene, also termed a selectable marker. Typical selection genes encode proteins that (a) confer resistance to antibioticsor other toxins, *e.g.*, ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies, or (c) supply critical nutrients not available from complex media, *e.g.*, the gene encoding D-alanine racemase for *Bacilli*. One example of a selection scheme utilizes a drug to arrest growth of a host cell. Those cells that are successfully transformed with a heterologous gene produce a protein conferring drug resistance and thus survive the selection regimen. Examples of such dominant selection use the drugs neomycin (Southern *et al.*, J. Molec. Appl. Genet. 1:327 (1982)), mycophenolic acid (Mulligan *et al.*, Science 209:1422 (1980)) or hygromycin (Sugden *et al.*, Mol. Cell. Biol. 5:410-413 (1985)). The three examples

given above employ bacterial genes under eukaryotic control to convey resistance to the appropriate drug G418 or neomycin (geneticin), xgpt (mycophenolic acid), or hygromycin, respectively.

[0101] Another example of suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up the heteromultimer nucleic acid, such as DHFR or thymidine kinase. The mammalian cell transformants are placed under selection pressure that only the transformants are uniquely adapted to survive by virtue of having taken up the marker. Selection pressure is imposed by culturing the transformants under conditions in which the concentration of selection agent in the medium is successively changed, thereby leading to amplification of both the selection gene and the DNA that encodes heteromultimer. Increased quantities of heteromultimer are synthesized from the amplified DNA. Other examples of amplifiable genes include metallothionein-I and -II, preferably primate metallothionein genes, adenosine deaminase, ornithine decarboxylase, etc.

5

10

15

20

25

30

40

45

50

55

[0102] For example, cells transformed with the DHFR selection gene are first identified by culturing all of the transformants in a culture medium that contains methotrexate (Mtx), a competitive antagonist of DHFR. An appropriate host cell when wild-type DHFR is employed is the Chinese hamster ovary (CHO) cell line deficient in DHFR activity, prepared and propagated as described by Urlaub and Chasin, Proc. Natl. Acad. Sci. USA 77:4216 (1980). The transformed cells are then exposed to increased levels of methotrexate. This leads to the synthesis of multiple copies of the DHFR gene, and, concomitantly, multiple copies of other DNA comprising the expression vectors, such as the DNA encoding the components of the heteromultimer. This amplification technique can be used with any otherwise suitable host, *e.g.*, ATCC No. CCL61 CHO-K1, notwithstanding the presence of endogenous DHFR if, for example, a mutant DHFR gene that is highly resistant to Mtx is employed (EP 117,060).

[0103] Alternatively, host cells (particularly wild-type hosts that contain endogenous DHFR) transformed or co-transformed with DNA sequences encoding heteromultimer, wild-type DHFR protein, and another selectable marker such as aminoglycoside 3'-phosphotransferase (APH) can be selected by cell growth in medium containing a selection agent for the selectable marker such as an aminoglycosidic antibiotic, *e.g.*, kanamycin, neomycin, or G418. See U.S. Patent No. 4,965,199.

[0104] A suitable selection gene for use in yeast is the *trp*1 gene present in the yeast plasmid YRp7 (Stinchcomb *et al.*, Nature 282:39 (1979); Kingsman *et al.*, Gene 7:141 (1979); or Tschemper *et al.*, Gene 10:157 (1980)). The *trp*1 gene provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, for example, ATCC No. 44076 or PEP4-1 (Jones, Genetics 85:12 (1977)). The presence of the *trp*1 lesion in the yeast host cell genome then provides an effective environment for detecting transformation by growth in the absence of tryptophan. Similarly, *Leu*2-deficient yeast strains (ATCC 20,622 or 38,626) are complemented by known plasmids bearing the *Leu*2 gene.

[0105] In addition, vectors derived from the 1.6 µm circular plasmid pKD 1 can be used for transformation of *Kluy-veromyces* yeasts. Bianchi *et al.*, Curr. Genet. 12:185 (1987). More recently, an expression system for large-scale production of recombinant calf chymosin was reported for *K. lactis*. Van den Berg, Bio/Technology 8:135 (1990). Stable multi-copy expression vectors for secretion of mature recombinant human serum albumin by industrial strains of *Kluy-veromyces* have also been disclosed (Fleer *et al.*, Bio/Technology 9:968-975 (1991)).

[0106] Expression and cloning vectors usually contain a promoter that is recognized by the host organism and is operably linked to the heteromultimer nucleic acid. A large number of promoters recognized by a variety of potential host cells are well known. These promoters are operably linked to heteromultimer-encoding DNA by removing the promoter from the source DNA by restriction enzyme digestion and inserting the isolated promoter sequence into the vector.

[0107] Promoters suitable for use with prokaryotic hosts include the β-lactamase and lactose promoter systems (Chang *et al.*, Nature 275:615 (1978); and Goeddel *et al.*, Nature 281:544 (1979)), alkaline phosphatase, a tryptophan (trp) promoter system (Goeddel, Nucleic Acids Res., 8:4057 (1980) and EP 36,776) and hybrid promoters such as the tac promoter (deBoer *et al.*, Proc. Natl. Acad. Sci. USA 80:21-25 (1983)). However, other known bacterial promoters are suitable. Their nucleotide sequences have been published, thereby enabling a skilled worker operably to ligate them to DNA encoding the heteromultimer (Siebenlist *et al.*, Cell 20:269 (1980)) using linkers or adaptors to supply any required restriction sites. Promoters for use in bacterial systems also will contain a Shine-Dalgamo(S.D.) sequence operably linked to the DNA encoding the heteromultimer.

[0108] Promoter sequences are known for eukaryotes. Virtually all eukaryotic genes have an AT-rich region located approximately25 to 30 bases upstream from the site where transcription is initiated. Another sequence found 70 to 80 bases upstream from the start of transcription of many genes is a CXCAAT region where X may be any nucleotide. At the 3' end of most eukaryotic genes is an AATAAA sequence that may be the signal for addition of the poly A tail to the 3' end of the coding sequence. All of these sequences are suitably inserted into eukaryotic expression vectors.

[0109] Examples of suitable promoting sequences for use with yeast hosts include the promoters for 3-phosphoglyceratekinase (Hitzeman *et al.*, J. Biol. Chem. <u>255</u>:2073 (1980)) or other glycolytic enzymes (Hess *et al.*, J. Adv. Enzyme Reg. <u>7</u>:149 (1968); and Holland, Biochemistry <u>17</u>:4900 (1978)), such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphofructokinase, glyceraldehydrogenase, become and glyceraldehydrogenase.

phoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase.

[0110] Other yeast promoters, which are inducible promoters having the additional advantage of transcription controlled by growth conditions, are the promoterregions for alcohol dehydrogenase2, isocytochrome C, acid phosphatase, degradative enzymes associated with nitrogen metabolism, metallothionein, glyceraldehyde-3-phosphate dehydrogenase, and enzymes responsible for maltose and galactose utilization. Suitable vectors and promoters for use in yeast expression are further described in Hitzeman *et al.*, EP 73,657A. Yeast enhancers also are advantageously used with yeast promoters.

5

10

20

25

30

40

45

50

[0111] Heteromultimer transcription from vectors in mammalian host cells is controlled, for example, by promoters obtained from the genomes of viruses such as polyoma virus, fowlpox virus (UK 2,211,504 published 5 July 1989), adenovirus (such as Adenovirus 2), bovine papilloma virus, avian sarcoma virus, cytomegalovirus, a retrovirus, hepatitis-B virus and most preferably Simian Virus 40 (SV40), from heterologous mammalian promoters, *e.g.*, the actin promoter or an immunoglobulin promoter or from heat-shock promoters.

[0112] The early and late promoters of the SV40 virus are conveniently obtained as an SV40 restriction fragment that also contains the SV40 viral origin of replication. Fiers *et al.*, Nature $\underline{273}$:113 (1978); Mulligan and Berg, Science $\underline{209}$:1422-1427 (1980); Pavlakis *et al.*, Proc. Natl. Acad. Sci. USA $\underline{78}$:7398-7402 (1981). The immediate early promoter of the human cytomegalovirus is conveniently obtained as a *Hin*dIII E restriction fragment. Greenaway *et al.*, Gene $\underline{18}$: 355-360 (1982). A system for expressing DNA in mammalian hosts using the bovine papilloma virus as a vector is disclosed in U.S. Patent No. 4,419,446. A modification of this system is described in U.S. Patent No. 4,601,978. See also Gray *et al.*, Nature $\underline{295}$:503-508 (1982) on expressing cDNA encoding immune interferon in monkey cells; Reyes *et al.*, Nature $\underline{297}$:598-601 (1982) on expression of human β -interferon cDNA in mouse cells under the control of a thymidine kinase promoter from herpes simplex virus; Canaani and Berg, Proc. Natl. Acad. Sci. USA $\underline{79}$:5166-5170 (1982) on expression of the human interferon β 1 gene in cultured mouse and rabbit cells; and Gorman *et al.*, Proc. Natl. Acad. Sci. USA $\underline{79}$:6777-6781 (1982) on expression of bacterial CAT sequences in CV-1 monkey kidney cells, chicken embryo fibroblasts, Chinese hamster ovary cells, HeLa cells, and mouse NIH-3T3 cells using the Rous sarcoma virus long terminal repeat as a promoter.

[0113] Transcription of DNA encoding the heteromultimer components by higher eukaryotes is often increased by inserting an enhancer sequence into the vector. Enhancers are relatively orientation and position independent, having been found 5' (Laimins *et al.*, Proc. Natl. Acad. Sci. USA 78:993 (1981)) and 3' (Lusky *et al.*, Mol. Cell Bio. 3:1108 (1983)) to the transcription unit, within an intron (Banerji *et al.*, Cell 3:1293 (1983)), as well as within the coding sequence itself (Osborne *et al.*, Mol. Cell Bio. 4:1293 (1984)). Many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, α -fetoprotein, and insulin). Typically, however, one will use an enhancer from a eukaryotic cell virus. Examples include the SV40 enhancer on the late side of the replication origin (bp 100-270), the cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers. See also Yaniv, Nature 2:97:17-18 (1982) on enhancing elements for activation of eukaryotic promoters. The enhancer may be spliced into the vector at a position 5' or 3' to the heteromultimer-encodingsequence, but is preferably located at a site 5' from the promoter.

[0114] Expression vectors used in eukaryotic host cells (yeast, fungi, insect, plant, animal, human, or nucleated cells from other multicellularorganisms) will also contain sequences necessary for the termination of transcription and for stabilizing the mRNA. Such sequences are commonly available from the 5' and, occasionally 3', untranslated regions of eukaryoticor viral DNAs or cDNAs. These regions contain nucleotide segments transcribed as polyadenylated fragments in the untranslated portion of the mRNA encoding the heteromultimer.

[0115] Construction of suitable vectors containing one or more of the above listed components employs standard ligation techniques. Isolated plasmids or DNA fragments are cleaved, tailored, and religated in the form desired to generate the plasmids required.

[0116] For analysis to confirm correct sequences in plasmids constructed, the ligation mixtures are used to transform *E. coli* K12 strain 294 (ATCC 31,446) and successful transformants selected by ampicillin or tetracycline resistance where appropriate. Plasmids from the transformants are prepared, analyzed by restriction endonuclease digestion, and/or sequenced by the method of Messing *et al.*, Nucleic Acids Res. 9:309 (1981) or by the method of Maxam *et al.*, Methods in Enzymology 65:499 (1980).

[0117] Particularly useful in the practice of this invention are expression vectors that provide for the transient expression in mammalian cells of DNA encoding heteromultimer. In general, transient expression involves the use of an expression vector that is able to replicate efficiently in a host cell, such that the host cell accumulates many copies of the expression vector and, in turn, synthesizes high levels of a desired polypeptide encoded by the expression vector. Sambrook *et al.*, *supra*, pp. 16.17 - 16.22. Transient expression systems, comprising a suitable expression vector and a host cell, allow for the convenient positive identification of polypeptidesencoded by cloned DNAs, as well as for the rapid screening of heteromultimers having desired binding specificities/affinities or the desired gel migration characteristics relative to heteromultimers or homomultimers lacking the non-natural disulfide bonds generated according to the instant invention.

[0118] Other methods, vectors, and host cells suitable for adaptation to the synthesis of the heteromultimer in recombinant vertebrate cell culture are described in Gething *et al.*, Nature 293:620-625 (1981); Mantei *et al.*, Nature 281: 40-46 (1979); EP 117,060; and EP 117,058. A particularly useful plasmid for mammalian cell culture expression of the heteromultimer is pRK5 (EP 307,247) or pSVI6B (PCT pub. no. WO 91/08291 published 13 June 1991).

5

10

20

25

30

40

45

50

[0119] The choice of host cell line for the expression of heteromultimer depends mainly on the expression vector. Another consideration is the amount of protein that is required. Milligram quantities often can be produced by transient transfections. For example, the adenovirus EIA-transformed 293 human embryonic kidney cell line can be transfected transiently with pRK5-based vectors by a modification of the calcium phosphate method to allow efficient heteromultimerexpression. CDM8-basedvectors can be used to transfect COS cells by the DEAE-dextran method (Aruffo *et al.*, Cell 61:1303-1313 (1990); and Zettmeissl *et al.*, DNA Cell Biol. (US) 9:347-353 (1990)). If larger amounts of protein are desired, the immunoadhesin can be expressed after stable transfection of a host cell line. For example, a pRK5-based vector can be introduced into Chinese hamster ovary (CHO) cells in the presence of an additional plasmid encoding dihydrofolate reductase (DHFR) and conferring resistance to G418. Clones resistant to G418 can be selected in culture. These clones are grown in the presence of increasing levels of DHFR inhibitor methotrexate and clones are selected in which the number of gene copies encoding the DHFR and heteromultimer sequences is co-amplified. If the immunoadhesin contains a hydrophobic leader sequence at its N-terminus, it is likely to be processed and secreted by the transfected cells. The expression of immunoadhesins with more complex structuresmay require uniquely suited host cells. For example, components such as light chain or J chain may be provided by certain myeloma or hybridoma host cells (Gascoigne *et al.*, *supra*; and Martin *et al.*, J. Virol. 67:3561-3568 (1993)).

[0120] Other suitable host cells for cloning or expressing the vectors herein are prokaryote, yeast, or other higher eukaryote cells described above. Suitable prokaryotes for this purpose include eubacteria, such as Gram-negative or Gram-positiveorganisms, for example, Enterobacteriaceaesuch as *Escherichia*, *e.g.*, *E. coli*, *Enterobacter*, *Erwinia*, *Klebsiella*, *Proteus*, *Salmonella*, *e.g.*, *Salmonella typhimurium*, *Serratia*, *e.g.*, *Serratia marcescans*, and *Shigella*, as well as *Bacilli* such as *B. subtilis* and *B. licheniformis* (*e.g.*, *B. licheniformis* 41P disclosed in DD 266,710 published 12 April 1989), *Pseudomonas* such as *P. aeruginosa*, and *Streptomyces*. One preferred *E. coli* cloning host is *E. coli* 294 (ATCC 31,446), although other strains such as *E. coli* B, *E. coli* X 1776 (ATCC 31,537), and *E. coli* W3110 (ATCC 27,325) are suitable. These examples are illustrative rather than limiting. Strain W3110 is a particularly preferred host or parent host because it is a common host strain for recombinant DNA product fermentations. Preferably, the host cell should secrete minimal amounts of proteolytic enzymes. For example, strain W3110 may be modified to effect a genetic mutation in the genes encoding proteins, with examples of such hosts including *E. coli* W3110 strain 27C7. The complete genotype of 27C7 is *tonA*Δ *ptr3 phoA*Δ*E15* Δ(*argF-lac*)169 *ompT*Δ *degP41kan^f*. Strain 27C7 was deposited on 30 October 1991 in the American Type Culture Collection as ATCC No. 55,244. Alternatively, the strain of *E. coli* having mutant periplasmic protease disclosed in U.S. Patent No. 4,946,783 issued 7 August 1990 may be employed. Alternatively, methods of cloning, *e.g.*, PCR or other nucleic acid polymerase reactions, are suitable.

[0121] In addition to prokaryotes, eukaryoticmicrobes such as filamentous fungi or yeast are suitable cloning or expression hosts for heteromultimer-encoding vectors. *Saccharomyces cerevisiae*, or common baker's yeast, is the most commonly used among lower eukaryotic host microorganisms. However, a number of other genera, species, and strains are commonly available and useful herein, such as *Schizosaccharomyces pombe* (Beach and Nurse, Nature 290:140 (1981); EP 139,383 published May 2, 1985); *Kluyveromyces* hosts (U.S. Patent No. 4,943,529; *Fleer et al.*, *supra*) such as, *e.g.*, *K. lactis* (MW98-8C, CBS683, CBS4574; Louvencourt *et al.*, J. Bacteriol., 737 (1983)), *K. fragilis* (ATCC 12,424), *K. bulgaricus* (ATCC 16,045), *K. wickeramii* (ATCC 24,178), *K. waltii* (ATCC 56,500), *K. drosophilarum* (ATCC 36,906; Van den Berg *et al.*, *supra*), *K thermotolerans*, and *K. marxianus*; *yarrowia* (EP 402,226); *Pichia pastoris* (EP 183,070; Sreekrishna *et al.*, J. Basic Microbiol. 28:265-278 (1988)); *Candida; Trichodermareesia* (EP 244,234); *Neurosporacrassa* (Case *et al.*, Proc. Natl. Acad. Sci. USA 76:5259-5263 (1979)); *Schwanniomyces* such as *Schwanniomyces occidentalis* (EP 394,538 published 31 October 1990); and filamentous fungi such as, *e.g.*, *Neurospora, Penicillium, Tolypocladium* (WO 91/003 57 published 10 January 1991), and *Aspergillus* hosts such as *A. nidulans* (Ballance *et al.*, Biochem. Biophys. Res. Commun. 112:284-289 (1983); Tilburn *et al.*, Gene 26:205-221 (1983); Yelton *et al.*, Proc. Natl. Acad. Sci. USA 81:1470-1474 (1984)) and *A. niger* (Kelly and Hynes, EMBO J. 4:475-479 (1985)).

Such host cells are capable of complex processing and glycosylation activities. In principle, any higher eukaryotic cell culture is workable, whether from vertebrate or invertebrate culture. Examples of invertebratecells include plant and insect cells. Numerous baculoviral strains and variants and corresponding permissive insect host cells from hosts such as *Spodopterafrugiperda* (caterpillar), *Aedes aegypti* (mosquito), *Aedes albopictus* (mosquito), *Drosophila melanogaster* (fruitfly), and *Bombyx mori* have been identified. See, *e.g.*, Luckow *et al.*, Bio/Technology 6:47-55 (1988); Miller *et al.*, in Genetic Engineering, Setlow *et al.*, eds., Vol. 8 (Plenum Publishing, 1986), pp. 277-279; and *Maeda et al.*, Nature 315:592-594 (1985). A variety of viral strains for transfection are publicly available, *e.g.*, the L-1 variant of *Autographa californica* NPV and the Bm-5 strain of *Bombyx mori* NPV, and such viruses may be used as the virus herein according to the present invention, particularly for transfection of *Spodoptera frugiperda* cells.

[0123] Plant cell cultures of cotton, corn, potato, soybean, petunia, tomato, and tobacco can be utilized as hosts. Typical ly, plant cells are transfected by incubation with certain strains of the bacterium *Agrobacterium tumefaciens*, which has been previously manipulated to contain the heteromultimer DNA. During incubation of the plant cell culture with *A. tumefaciens*, the DNA encoding the heteromultimer is transferred to the plant cell host such that it is transfected, and will, under appropriate conditions, express the heteromultimer DNA. In addition, regulatory and signal sequences compatible with plant cells are available, such as the nopaline synthase promoter and polyadenylation signal sequences. Depicker *et al.*, J. Mol. Appl. Gen. 1:561 (1982). In addition, DNA segments isolated from the upstream region of the T-DNA *780* gene are capable of activating or increasing transcription levels of plant-expressible genes in recombinant DNA-containing plant tissue. EP 321, 196 published 21 June 1989.

5

10

15

20

25

30

40

45

50

55

[0124] The preferred hosts are vertebrate cells, and propagation of vertebrate cells in culture (tissue culture) has become a routine procedure in recent years (Tissue Culture, Academic Press, Kruse and Patterson, editors (1973)). Examples of useful mammalian host cell lines are monkey kidney CV1 line transformed by SV40 (COS-7, ATCC CRL 1651); human embryonic kidney line (293 or 293 cells subcloned for growth in suspension culture, Graham *et al.*, J. Gen Virol. 36:59 (1977)); baby hamster kidney ceils (BHK, ATCC CCL 10); Chinese hamster ovary cells/-DHFR (CHO, Urlaub and Chasin, Proc. Natl. Acad. Sci. USA 77:4216 (1980)); mouse sertoli cells (TM4, Mather, Biol. Reprod. 23: 243-251 (1980)); monkey kidney cells (CV1 ATCC CCL 70); African green monkey kidney cells (VERO-76, ATCC CRL-1587); human cervical carcinomacells (HELA, ATCC CCL 2); canine kidney cells (MDCK, ATCC CCL 34); buffalo rat liver cells (BRL 3A, ATCC CRL 1442); human lung cells (W138, ATCC CCL 75); human liver cells (Hep G2, HB 8065); mouse mammary tumor (MMT 060562, ATCC CCL51); TRI cells (Mather *et al.*, Annals N.Y. Acad. Sci. 383:44-68 (1982)); MRC 5 cells; FS4 cells; and a human hepatoma line (Hep G2).

[0125] Host cells are transfected with the above-described expression or cloning vectors of this invention and cultured in conventional nutrient media modified as appropriate for inducing promoters, selecting transformants, or amplifying the genes encoding the desired sequences. Depending on the host cell used, transfection is done using standard techniques appropriate to such cells. The calcium treatment employing calcium chloride, as described in section 1.82 of Sambrook *et al.*, *supra*, or electroporation is generally used for prokaryotes or other cells that contain substantial cell-wall barriers. Infection with *Agrobacterium tumefaciens* is used for transformation of certain plant cells, as described by Shaw *et al.*, Gene 23:315 (1983) and WO 89/05859 published 29 June 1989. In addition, plants may be transfected using ultrasound treatment as described in WO 91/00358 published 10 January 1991.

[0126] For mammalian cells without such cell walls, the calcium phosphate precipitationmethod of Graham and van der Eb, Virology 52:456-457 (1978) is preferred. General aspects of mammalian cell host system transformations have been described by Axel in U.S. Patent No. 4,399,216 issued 16 August 1983. Transformations into yeast are typically carried out according to the method of Van Solingen *et al.*, J. Bact. 130:946 (1977) and Hsiao *et al.*, Proc. Natl. Acad. Sci. (USA) 76:3829 (1979). However, other methods for introducing DNA into cells, such as by nuclear microinjection, electroporation, bacterial protoplast fusion with intact cells, or polycations, *e.g.*, polybrene, polyornithine, *etc.*, may also be used. For various techniques for transforming mammalian cells, see Keown *et al.*, Methods in Enzymology (1989), Keown *et al.*, Methods in Enzymology 185:527-537 (1990), and Mansour *et al.*, Nature 336:348-352 (1988).

[0127] Prokaryot ic cells used to produce the heteromultimer polypeptide of this invention are cultured in suitable media as described generally in Sambrook *et al.*, *supra*.

[0128] The mammalian host cells used to produce the heteromultimer of this invention may be cultured in a variety of media. Commercially available media such as Ham's F10 (Sigma), Minimal Essential Medium ((MEM), Sigma), RPMI-1640 (Sigma), and Dulbecco's Modified Eagle's Medium ((DMEM), Sigma) are suitable for culturing the host cells. In addition, any of the media described in Ham and Wallace, Meth. Enz. <u>58</u>:44 (1979), Barnes and Sato, Anal. Biochem. <u>102</u>:255 (1980), U.S. Patent Nos. 4,767,704; 4,657,866; 4,927,762; or 4,560,655; WO 90/03430; WO 87/00195; U.S. Patent Re. 30,985; or U.S. Patent No. 5,122,469, may be used as culture media for the host cells. Any of these media may be supplemented as necessary with hormones and/or other growth factors (such as insulin, transferrin, or epidermal growth factor), salts (such as sodium chloride, calcium, magnesium, and phosphate), buffers (such as HEPES), nucleosides (such as adenosine and thymidine), antibiotics (such as Gentamycin™ drug), trace elements (defined as inorganic compounds usually present at final concentrations in the micromolar range), and glucose or an equivalent energy source. Any other necessary supplements may also be included at appropriate concentrations that would be known to those skilled in the art. The culture conditions, such as temperature, pH, and the like, are those previously used with the host cell selected for expression, and will be apparent to the ordinarily skilled artisan.

[0129] In general, principles, protocols, and practical techniques for maximizing the productivity of mammalian cell cultures can be found in Mammalian Cell Biotechnology: a Practical Approach, M. Butler, ed., IRL Press, 1991.

[0130] The host cells referred to in this disclosure encompass cells in culture as well as cells that are within a host animal.

4. Recovery of the Heteromultimer

10

15

20

25

30

40

45

50

55

[0131] The heteromultimer preferably is generally recovered from the culture medium as a secreted polypeptide, although it also may be recovered from host cell lysate when directly produced without a secretory signal. If the heteromultimer is membrane-bound, it can be released from the membrane using a suitable detergent solution (*e.g.* Triton-X 100).

[0132] When the heteromultimer is produced in a recombinant cell other than one of human origin, it is completely free of proteins or polypeptides of human origin. However, it is necessary to purify the heteromultimer from recombinant cell proteins or polypeptides to obtain preparations that are substantially homogeneous as to heteromultimer. As a first step, the culture medium or lysate is normally centrifuged to remove particulate cell debris.

[0133] Heterodimers having antibody constant domains can be conveniently purified by hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography, with affinity chromatography being the preferred purification technique. Where the heteromultimer comprises a C_H3 domain, the Bakerbond ABX™ resin (J. T. Baker, Phillipsburg, NJ) is useful for purification. Other techniques for protein purification such as fractionation on an ion-exchange column, ethanol precipitation, reverse phase HPLC, chromatography on silica, chromatography on heparin Sepharose, chromatography on an anion or cation exchange resin (such as a polyaspartic acid column), chromatofocusing, SDS-PAGE, and ammonium sulfate precipitation are also available depending on the polypeptide to be recovered. The suitability of protein A as an affinity ligand depends on the species and isotype of the immunoglobulin Fc domain that is used in the chimera. Protein A can be used to purify immunoadhesinsthat are based on human γ 1, γ 2, or γ 4 heavy chains (Lindmarket al., J. Immunol. Meth. 62:1-13 (1983)). Protein G is recommended for all mouse isotypes and for human γ 3 (Guss et al., EMBO J. 5:15671575 (1986)). The matrix to which the affinity ligand is attached is most often agarose, but other matrices are available. Mechanically stable matrices such as controlled pore glass or poly(styrenedivinyl)benzene allow for faster flow rates and shorter processing times than can be achieved with agarose. The conditions for binding an immunoadhesin to the protein A or G affinity column are dictated entirely by the characteristics of the Fc domain; that is, its species and isotype. Generally, when the proper ligand is chosen, efficient binding occurs directly from unconditioned culture fluid. One distinguishing feature of immunoadhesins is that, for human γ 1 molecules, the binding capacity for protein A is somewhat diminished relative to an antibody of the same Fc type. Bound immunoadhesin can be efficiently eluted either at acidic pH (at or above 3.0), or in a neutral pH buffer containing a mildly chaotropic salt. This affinity chromatography step can result in a heterodimer preparation that is >95% pure.

5. Uses for a Heteromultimeric Multispecific Antibody Having Common Light Chains

Many therapeutic applications for the heteromultimer are contemplated. For example, the heteromultim er can be used for redirected cytotoxicity (e.g. to kill tumor cells), as a vaccine adjuvant, for delivering thrombolytic agents to clots, for converting enzyme activated prodrugs at a target site (e.g. a tumor), for treating infectious diseases, targeting immune complexes to cell surface receptors, or for delivering immunotoxinsto tumor cells. For example, tumor vasculature targeting has been accomplished by targeting a model endothelial antigen, class II major histocompatibility complex, with an antibody-ricin immunotoxin (Burrows, F.J. and Thorpe, P.E. (1993) Proc Natl Acad Sci USA 90: 8996-9000). Significantly greater efficacy was achieved by combining the anti-endothelial immunotoxin with a second immunotoxin directed against the tumor cells themselves (Burrows, F.J. and Thorpe, P.E. (1993) supra). Recently, tissue factor was successfully targeted to tumor vasculature using a bispecific antibody, triggering local thrombosis that resulted in significant anti-tumor efficacy (Huang, X. et al. (1997) Science 275:547-550). In addition, bispecific diabodies have been used successfully to direct cytotoxic T-cells to kill target breast tumor cells and B-cell lymphoma cells in vitro (Zhu, Z. et al. (1996) Bio/Technology 14:192-196; and Holliger, P. et al. (1996) Protein Engin. 9:299-305). [0135] Therapeutic formulations of the heteromultimer are prepared for storage by mixing the heteromultimer having the desired degree of purity with optional physiologically acceptable carriers, excipients, or stabilizers (Remington's Pharmaceutical Sciences, 16th edition, Osol, A., Ed., (1980)), in the form of lyophilized cake or aqueous solutions. Acceptable carriers, excipients or stabilizers are nontoxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate, and other organic acids; antioxidants including ascorbic acid; low molecular weight (less than about 10 residues) polypeptides; proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine, arginine or lysine; monosaccharides, disaccharides, and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; salt-forming counterions such as sodium; and/or nonionic surfactants such as Tween, Pluronics or polyethylene glycol (PEG).

[0136] The heteromultimer also may be entrapped in microcapsulesprepared, for example, by coacervation techniques or by interfacial polymerization (for example, hydroxymethylcellulose or gelatin-microcapsules and poly-[meth-ylmethacylate]microcapsules, respectively), in colloidal drug delivery systems (for example, liposomes, albumin microspheres, microemulsions, nano-particles and nanocapsules), or in macroemulsions. Such techniques are disclosed in

Remington's Pharmaceutical Sciences, supra.

5

10

15

20

25

30

40

45

50

55

[0137] The heteromultimerto be used for *in vivo* administrationmust be sterile. This is readily accomplished by filtration through sterile filtration membranes, prior to or following lyophilization and reconstitution. The heteromultimer ordinarily will be stored in lyophilized form or in solution.

[0138] Therapeutic heteromultimer compositions generally are placed into a containerhaving a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

[0139] The route of heteromultimer administration is in accord with known methods, *e.g.*, injection or infusion by intravenous, intraperitoneal, intracerebral, intramuscular, intraocular, intraarterial, or intralesional routes, or by sustained release systems as noted below. The heteromultimer is administered continuously by infusion or by bolus injection.

[0140] Suitable examples of sustained-release preparations include semipermeable matrices of solid hydrophobic polymers containing the protein, which matrices are in the form of shaped articles, *e.g.*, films, or microcapsules. Examples of sustained-release matrices include polyesters, hydrogels (*e.g.*, poly(2-hydroxyethyl-methacrylate) as described by Langer *et al.*, J. Biomed. Mater. Res. <u>15</u>:167-277 (1981) and Langer, Chem. Tech. <u>12</u>:98-105 (1982) or poly (vinylalcohol)), polylactides (U.S. Patent No. 3,773,919, EP 58,481), copolymers of L-glutamic acid and gamma ethyl-L-glutamate (Sidman *et al.*, Biopolymers <u>22</u>:547-556 (1983)), non-degradableethylene-vinylacetate (Langeret *al.*, *su-pra*), degradable lactic acid-glycolic acid copolymers such as the Lupron Depot™ (injectable microspheres composed of lactic acid-glycolic acid copolymer and leuprolide acetate), and poly-D-(-)-3-hydroxybutyric acid (EP 133,988).

[0141] While polymers such as ethylene-vinyl acetate and lactic acid-glycolic acid enable release of molecules for over 100 days, certain hydrogels release proteins for shorter time periods. When encapsulated proteins remain in the body for a long time, they may denature or aggregate as a result of exposure to moisture at 37°C, resulting in a loss of biological activity and possible changes in immunogenicity. Rational strategies can be devised for protein stabilization depending on the mechanism involved. For example, if the aggregation mechanism is discovered to be intermolecular S-S bond formation through thio-disulfide interchange, stabilization may be achieved by modifying sulfhydryl residues, lyophilizing from acidic solutions, controlling moisture content, using appropriate additives, and developing specific polymer matrix compositions.

[0142] Sustained-release heteromultimer compositions also include liposomally entrapped heteromultimer. Liposomes containing heteromultimer are prepared by methods known *per se:* DE 3,218,121; Epstein *et al.*, Proc. Natl. Acad. Sci. USA <u>82</u>:3688-3692 (1985); Hwang *et al.*, Proc. Natl. Acad. Sci. USA <u>77</u>:4030-4034 (1980); EP 52,322; EP 36,676; EP 88,046; EP 143,949; EP 142,641; Japanese patent application 83-118008; U.S. Patent Nos. 4,485,045 and 4,544,545; and EP 102,324. Ordinarily the liposomes are of the small (about 200-800 Angstroms) unilamellar type in which the lipid content is greater than about 30 mol. % cholesterol, the selected proportion being adjusted for the optimal heteromultimer therapy.

[0143] An effective amount of heteromultimer to be employed therapeutically will depend, for example, upon the therapeutic objectives, the route of administration, and the condition of the patient. Accordingly, it will be necessary for the therapist to titer the dosage and modify the route of administration as required to obtain the optimal therapeutic effect. A typical daily dosage might range from about 1 µg/kg to up to 10 mg/kg or more, depending on the factors mentioned above. Typically, the clinician will administer heteromultimer until a dosage is reached that achieves the desired effect. The progress of this therapy is easily monitored by conventional assays.

[0144] The heteromultimers described herein can also be used in enzyme immunoassays. To achieve this, one arm of the heteromultimer can be designed to bind to a specific epitope on the enzyme so that binding does not cause enzyme inhibition, the other arm of the heteromultimer can be designed to bind to the immobilizing matrix ensuring a high enzyme density at the desired site. Examples of such diagnostic heteromultimers include those having specificity for IgG as well as ferritin, and those having binding specificities for horse radish peroxidase (HRP) as well as a hormone, for example.

[0145] The heteromultimerscan be designed for use in two-site immunoassays. For example, two bispecific heteromultimers are produced binding to two separate epitopes on the analyte protein - one heteromultimer binds the complex to an insoluble matrix, the other binds an indicator enzyme.

[0146] Heteromultimers can also be used for *in vitro* or *in vivo* immunodiagnosis of various diseases such as cancer. To facilitate this diagnostic use, one arm of the heteromultimer can be designed to bind a tumor associated antigen and the other arm can bind a detectable marker (e.g. a chelatorwhich binds a radionuclide). For example, a heteromultimerhaving specificities for the tumor associated antigen CEA as well as a bivalent hapten can be used for imaging of colorectal and thryroid carcinomas. Other non-therapeutic, diagnostic uses for the heteromultimer will be apparent to the skilled practitioner.

[0147] For diagnostic applications, at least one arm of the heteromultimer typically will be labeled directly or indirectly with a detectable moiety. The detectable moiety can be any one which is capable of producing, either directly or indirectly, a detectable signal. For example, the detectable moiety may be a radioisotope, such as ³H, ¹⁴C, ³²P, ³⁵S, or ¹²⁵I; a fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or luciferin; or an en-

zyme, such as alkaline phosphatase, beta-galactosidase or horseradish peroxidase (HRP).

[0148] Any method known in the art for separately conjugating the heteromultimer to the detectable moiety may be employed, including those methods described by Hunter *et al.*, Nature 144:945 (1962); David *et al.*, Biochemistry 13: 1014 (1974); Pain *et al.*, J. Immunol. Meth. 40:219 (1981); and Nygren, J. Histochem. and Cytochem. 30:407 (1982).

[0149] The heteromultimers of the present invention may be employed in any known assay method, such as competitive binding assays, direct and indirect sandwich assays, and immunoprecipitation assays. Zola, Monoclonal Antibodies: A Manual of Techniques, pp.147-158 (CRC Press, Inc., 1987).

[0150] Competitive binding assays rely on the ability of a labeled standard to compete with the test sample analyte for binding with a limited amount of heteromultimer. The amount of analyte in the test sample is inversely proportional to the amount of standard that becomes bound to the heteromultimer. To facilitate determining the amount of standard that becomes bound, the heteromultimers generally are insolubilized before or after the competition, so that the standard and analyte that are bound to the heteromultimers may conveniently be separated from the standard and analyte which remain unbound.

[0151] The heteromu Itimers are particularly useful for sandwich assays which involve the use of two molecules, each capable of binding to a different immunogenic portion, or epitope, of the sample to be detected. In a sandwich assay, the test sample analyte is bound by a first arm of the heteromultimer which is immobilized a solid support, and thereafter a second arm of the heteromultimer binds to the analyte, thus forming an insoluble three part complex. See, e.g., US Pat No. 4,376,110. The second arm of the heteromultimermay itself be labeled with a detectable moiety (direct sandwich assays) or may be measured using an anti-immunoglobulinantibody that is labeled with a detectablemoiety (indirect sandwich assay). For example, one type of sandwich assay is an ELISA assay, in which case the detectable moiety is an enzyme.

[0152] Below are examples of specific embodiments for carrying out the present invention. The examples are offered for illustrative purposes only, and are not intended to limit the scope of the present invention in any way.

[0153] All publications, patents and patent applications cited herein, whether *supra* or *infra*, are hereby incorporated by reference in their entirety.

EXAMPLES

5

10

15

20

25

30

40

45

[0154] A strategy is presented for preparing Fc-containing BsAb (Fig. IC). In this strategy, we have engineered the C_H3 domain of antibody heavy chains so that they heterodimerize but do not homodimerize. This was accomplished by installing inter-chain disulfide bonds in the C_H3 domain in conjunction with sterically complimentary mutations obtained by rational design (Ridgway et al., *supra* (1996)) and phage display selection as described herein. Use of a single light chain for both antigen binding specificities circumvents the problem of light chain mispairing (Fig 1A-1C). Antibodies with the same light chain were readily isolated by panning a very large human scFv library (Vaughan, T. J., *et al.*, (1996) *supra*).

Example 1: Generation of protuberance-into-cavity heteromultimer immunoadhesins

[0155] The C_H3 interface between the humanized anti-CD3/CD4-lgG chimera previously described by Chamow *et al.* J. Immunol. 153:4268 (1994) was engineered to maximize the percentage of heteromultimers which could be recovered. Protuberance-into-cavityand wild-type C_H3 variants were compared in their ability to direct the formation of a humanized antibody-immunoadhesin chimera (Ab/la) anti-CD3/CD4-lgG.

[0156] Thus, mutations were constructed in the C_H3 domain of the humanized anti-CD3 antibody heavy chain and in CD4-lgG by site-directed mutagenesis using mismatched oligonucleotides (Kunkel *et al.*, Methods Enzymol. <u>154</u>: 367 (1987) and P. Carter, in Mutagenesis: a Practical Approach, M. J. McPherson, Ed., IRL Press, Oxford, UK, pp. 1-25 (1991)) and verified by dideoxynucleotide sequencing (Sanger *et al.*, Proc. Natl. Acad. Sci. USA <u>74</u>:5463 (1977)). See Table 3 below.

50

55

TABLE 3

Most Preferred Mutants C_H3 of anti-CD3 C_H3 of CD4-IgG Y407T T366Y Y407A T366W T394W F405A Y407T T366Y T366Y:F405A T394W:Y407T T366W:F405W T394S:Y407A F405W:Y407A T366W:T394S

	Preferred Mutants	
F405W	T394S	

25

30

45

50

55

5

10

15

20

[0157] Residue T366 is within hydrogen-bonding distance of residue Y407 on the partner C_H3 domain. Indeed the principal intermolecular contact to residue T366 is to residue Y407 and *vice versa*. One protuberance-into-cavitypair was created by inverting these residues with the reciprocal mutations of T366Y in one C_H3 domain and Y407T in the partner domain thus maintaining the volume of side chains at the interface. Mutations are denoted by the wild-type residue followed by the position using the Kabat numbering system (Kabat *et al.* (1991) *supra*) and then the replacement residue in single-letter code. Multiple mutations are denoted by listing component single mutations separated by a colon.

[0158] Phagemids encoding anti-CD3 light (L) and heavy (H) chain variants (Shalaby *et al.*, J. Exp. Med. <u>175</u>:217 (1992) and *Rodrigues et al.*, Int. J. Cancer (Suppi.) <u>7</u>:45 (1992)) were co-transfected into human embryonic kidney cells, 293S, together with a CD4-lgG variant encoding phagemid (Byrn *et al.*, Nature 344:667 (1990)) as previously described (Chamow *et al.*, J. Immunol. <u>153</u>:4268 (1994)). The total amount of transfectedphagemid DNAs was fixed whereas the ratio of different DNAs was varied to maximize the yield of Ab/la chimera. The ratio (by mass) of la: heavy chain: light chain input DNAs (15 µg total) was varied as follows: 8:1:3; 7:1:3; 6:1:3; 5:1:3; 4:1:3; 3:1:3; 1:0:0; 0:1:3.

[0159] The products were affinity purified using Staphylococcal protein A (ProSep A, BioProcessing Ltd, UK) prior to analysis by SDS-PAGE followed by scanning LASER densitometry. Excess light over heavy chain DNA was used to avoid the light chain from being limiting. The identity of products was verified by electroblotting on to PVDF membrane (Matsudaira, J. Biol. Chem. 262:10035 (1987)) followed by amino terminal sequencing.

[0160] Co-transfection of phagemids for light chain together with those for heavy chain and la incorporating wild-type C_H3 resulted in a mixture of Ab/la chimera, IgG and la homodimer products as expected (Chamow *et al.*, J. Immunol. <u>153</u>:4268(1994)). The larger the fraction of input DNA encoding antibody heavy plus light chains or 1a the higher the fraction of correspondinghomodimers recovered. An input DNA ratio of 6:1:3 of la:H:L yielded 54.5 % Ab/la chimera with similar fractions of 1a homodimer (22.5 %) and IgG (23.0 %). These ratios are in good agreement with those expected from equimolar expression of each chain followed by random assortment of heavy chains with no bias being introduced by the method of analysis: 50 % Ab/la chimera, 25 % la homodimer and 25 % IgG.

[0161] In contrast to chains containing wild-type C_H3, Ab/la chimera was recovered in yields of up to 92 % from cotransfections in which the anti-CD3 heavy chain and CD4-lgG 1a contained the Y407T cavity and T366Y protuberance mutations, respectively. Similar yields of antibody/immunoadhesin chimera were obtained if these reciprocal mutations were installed with the protuberance on the heavy chain and the cavity in the la. In both cases monomer was observed for the chain containing the protuberance but not the cavity. Without being limited to any one theory, it is believed that the T366Y protuberance is more disruptive to homodimer formation than the Y407T cavity. The fraction of Ab/la hybrid was not significantly changed by increasing the size of both protuberance and cavity (Ab T366W, la Y407A). A second protuberance and cavity pair (Ab F405A, la T394W) yielded up to 71 % Ab/la chimera using a small fraction of la input DNA to offset the unanticipated proclivity of the la T394W protuberance variant to homodimerize. Combining the two

independent protuberance-into-cavity mutant pairs (Ab T366Y:F405A, la T394W:Y407T) did not improve the yield of Ab/la hybrid over the Ab T366Y, la Y407T pair.

[0162] The fraction of Ab/la chimera obtained with T366Y and Y407T mutant pair was virtually independent of the ratio of input DNAs over the range tested. Furthermore the contaminating species were readily removed from the Ab/la chimera by ion exchange chromatography (0-300 mM NaCl in 20 mM Tris-HCl, pH8.0) on a mono S HR 5/5 column (Pharmacia, Piscataway, NJ). This augurs well for the preparation of larger quantities Ab/la chimeras using stable cell lines where the relative expression levels of Ab and la are less readily manipulated than in the transient expression system.

[0163] The protuberance-into-cavitymutations identified are anticipated increase the potential applications of Fccontaining BsAb by reducing the complexity of the mixture of products obtained from a possible ten major species (Suresh *et al.*, Methods Enzymol. 121:210 (1990)) down to four or less (Figs. 1A-1B). It is expected that the T366Y and Y407T mutant pair will be useful for generating heteromultimers of other human IgG isotypes (such as IgG₂, IgG₃ or IgG₄) since T366 and Y407 are fully conserved and other residues at the C_H3 domain interface of IgG₁ are highly conserved.

Example 2: Generation of non-naturally occurring disulfide linkages in heteromultimeric immunoadhesins

A. Design of C_H3 inter-chain disulfide bonds.

5

10

15

30

35

40

45

50

55

[0164] Three criteria were used to identify pairs of residues for engineering disulfide bond between partner C_H3 domains: i) The Cα separation preferably is similar to those found in natural disulfide bonds (5.0 to 6.8 Å) (Srinivasan, N., et al., Int. J. Peptides Protein Res. 36:147-155 (1990)). Distances of up to 7.6 Å were permitted to allow for main chain movement and to take into account the uncertainty of atomic positions in the low resolution crystal structure (Deisenhofer, Biochemistry 20:2361-2370 (1981)). ii) The Cα atoms should be on different residues on the two C_H3 domains. iii) The residues are positioned to permit disulfide bonding (Srinivasan, N., et al., (1990) supra).

[0165] B. Modeling of disulfide bonds. Disulfide bonds were modeled into the human IgG₁ Fc (Deisenhofer, *supra*) as described for humAb4D5-Fv (Rodrigues et al., Cancer Res. <u>55</u>:63-70 (1995)) using Insight II release 95.0 (Biosym/MSI).

[0166] C. Construction of C_H3 variants. Mutations were introduced into the C 3_H domain of a humanized anti-CD3 heavy chain or CD4-lgG by site-directed mutagenesis (Kunkel, *et al.*, Methods Enzymol. <u>154</u>:367-382 (1987)) using the following synthetic oligonucleotides:

Y349C, 5' CTCTTCCCGAGATGGGGGCAGGGTGCACACCTGTGG 3' (SEQ. ID NO: 1)

S354C, 5' CTCTTCCCGACATGGGGGCAG 3' (SEQ. ID NO: 2)

E356C, 5' GGTCATCTCACACCGGGATGG 3' (SEQ. ID NO: 3)

E357C, 5' CTTGGTCATACATTCACGGGATGG 3' (SEQ. ID NO: 4)

L351C, 5' CTCTTCCCGAGATGGGGGACAGGTGTACAC 3' (SEQ. ID NO: 5)

D399C, 5' GCCGTCGGAACACAGCACGGG 3' (SEQ. ID NO: 6)

K392C, 5' CTGGGAGTCTAGAACGGGAGGCGTGGTACAGTAGTTGTT 3' (SEQ. ID NO: 7)

T394C, 5' GTCGGAGTCTAGAACGGGAGGACAGGTCTTGTA 3' (SEQ. ID NO: 8)

V397C, 5' GTCGGAGTCTAGACAGGGAGG 3' (SEQ. ID NO: 9)

D399S, 5' GCCGTCGGAGCTCAGCACGGG 3' (SEQ. ID NO: 10)

K392S, 5' GGGAGGCGTGGTGCTGTAGTTGTT 3' (SEQ. ID NO: 11)

C231S:C234S 5'GTTCAGGTGCTGGGCTCGGTGGGCTTGTGTGAGTTTTG 3' (SEQ. ID NO: 12)

[0167] Mutations are denoted by the amino acid residue and number (Eu numbering scheme of Kabat *et al.*, *supra* (1991), followed by the replacement amino acid. Multiple mutations are represented by the single mutation separated by a colon. Mutants were verified by dideoxynucleotide sequencing (Sanger *et al.*, *supra* (1977)) using Sequenase version 2.0 (United States Biochemicals, Cleveland, OH).

[0168] D. An inter-chain disulfide enhances heterodimer formation. Six pairs of molecules containing inter-chain disulfide bonds in the C_H3 domain ("disulfide-C_H3" variants; v1-v6, Table 4) were compared with parent molecules in their ability to direct the formation of an Ab/la hybrid, anti-CD3/CD4-lgG (Chamow *et al.*, *supra* (1994)). Plasmids encoding CD4-lgG and anti-CD3 heavy chain variants were co-transfected into 293S cells, along with an excess of plasmid encoding the anti-CD3 light chain. The yield of heterodimer was optimized by transfecting with a range of la: H chain:L chain DNA ratios. The Ab/la heterodimer, lgG and la homodimer products were affinity-purifiedusing Staphylococcal protein A and quantified by SDS-PAGE and scanning laser densitometry (Ridgway *et al.*, *supra* (1996)).

[0169] Each disulfide-C_H3 pair gave rise to three major species, similar to the parent molecules. However, Ab/la heterodimer from disulfide-C_H3 variants was shifted in electrophoretic mobility, consistent with formation of an interchain disulfide in the C_H3 domain. Further evidence of disulfide bond formation was provided by the inter-chain disulfides in the hinge. Covalently bonded Ab/la hybrids were observed by SDS-PAGE for disulfide-C_H3 variants but not for molecules with wildtype C_H3 domains in which hinge cysteines were mutated to serine. Disulfide-C_H3 variants were prepared and designated Y349C/S354'C, Y349C/E356'C, Y349C/E357'C, L351C/E354'C, T394C/E397'C, and D399C/K392C. Only one variant (D399C/K392'C) substantially increased the yield of Ab/la hybrid over wildtype (76% vs. 52%, respectively) as determined by SDS-PAGE analysis of the variants. Mutations are denoted by the amino acid residue and number (Eu numbering scheme of Kabat et al. (1991) *supra*), followed by the replacement amino acid. Mutations in the first and second copies of C_H3 come before and after the slash, respectively. Residues in the second copy of C_H3 are designated with a prime ('). This improvement apparently reflects disulfide bond formation rather than replacement of residues K392 and D399, since the mutations K392S/D399'S gave both a similar Ab/la yield and Ab/la electrophoretic mobility relative to wildtype. Homodimers migrated similarly to those with wildtype Fc domains, demonstrating referential engineered inter-chain disulfide bond formation in the C_H3 domain of heterodimers. All disulfide-C_H3 variants were expressed at approximately the same level as the parent molecules in 293S cells.

[0170] E. Disulfides combined with protuberance-into-cavity engineering increases the yield of heterodimer to 95%. The best disulfide pair increased the percent of heterodimer to 76% and the protuberance-into-cavitystrategy increased the percent of heterodimer to 87% (Table 4; see also Ridgway *et al.*, (1996) *supra*). These two strategies rely on different principles to increase the probability of generating heterodim er. Therefore, we combined the two strategies, anticipating further improvement in the yield of heterodimer. Two of the modeled disulfides, containing L351C or T394C, could potentially form disulfide-bonded homodimers as well as disulfide-bonded heterodimers (L351C/S354'C and T394CN397'C), thus decreasing their utility. The remaining four disulfide pairs were installed into the phage-selected heterodimer (variants v9-v16) and assayed for the yield of heterodimer (Table 4). Yields of approximately 95% heterodimer were obtained. Again, the heterodimer showed an electrophoretic mobility shift compared to wildtype and v8 variants.

Table 4

	Yield	s of Heterodimers from C _H 3 Vari	ants
		Mutations	
Variant	Subunit A	Subunit B	Yield of heterodimer (%)
wildtype	-	-	51 ± 1
v1	Y349C	S354C	54 ± 4
v2	Y349C	E356C	55 ± 6
v3	Y349C	E357C	57±4
v4	L351C	E354C	56 ± 3

50

45

40

5

10

20

25

30

55

Table 4 (continued)

5

10

15

20

25

30

35

40

45

50

55

	Yield	s of Heterodimers from C _H 3 Vari	ants
		Mutations	
Variant	Subunit A	Subunit B	Yield of heterodimer (%)
v 5	T394C	E397C	57 ± 2
v6	D399C	K392C	73 ± 3
v7	D399S	L392S	55 ± 1
v8	T366W	T366S:L368A:Y407V	86.7 ± 2.3
v9	T366W: D399C	T366S:L368A: K392C :Y407V	86.5 ± 0.5
v11	S354C :T366W	Y349C :T366S:L368A:Y407V	95 ± 2
v12	E356C :T366W	Y349C :T366S:L368A:Y407V	94 ± 2
v13	E357C :T366W	Y349C :T366S:L368A:Y407V	93 ± 2
v14	T366W: K392C	T366S: D399C :L368A:Y407V	92 ± 1
v15	Y349C :T366W	S354C :T366S:L368A:Y407V	90 ± 1
v16	Y349C :T366W	E356C :T366S:L368A:Y407V	95.5 ±0.5
v17	Y349C :T366W	E357C :T366S:L368A:Y407V	91.0 ± 1.0

Example 3: Structure-guided phage display selection for complementary mutations that enhance protein-protein interaction in heteromultimers

[0171] The following strategy is useful in the selection of complementary mutations in polypeptides that interact at an interface via a multimerization domain. The strategy is illustrated below as it applies to the selection of complementary protuberance-into-cavity mutations. However, the example is not meant to be limiting and the strategy may be similarly applied to the selection of mutations appropriate for the formation of non-naturally occurring disulfide bonds, leucine zipper motifs, hydrophobic interactions, hydrophilic interactions, and the like.

A. <u>Phage display selection</u>. A phage display strategy was developed for the selection of stable C_H3 heterodimers and is diagramed in Fig.2. The selection uses a protuberance mutant, T366W (Ridgway *et al.*, *supra* (1996)), fused to a peptide flag (gD peptide flag, for example, Lasky, L. A. and Dowbenko, D. J. (1984) DNA <u>3</u>:23-29; and Berman, P. W., *et al.* (1985) Science <u>227</u>:1490-1492) that is coexpressed with a second copy of C_H3 fused to M13 gene III protein. A library of cavity mutants was created in this second copy of C_H3 by randomization of the closest neighboring residues to the protuberance on the first C_H3 domain. Phage displaying stable C_H3 heterodimers were then captured using an anti-flag Ab.

A C_H3 phage display library of 1.1 x 10^5 independent clones was constructed by replacement of a segment of the natural C_H3 gene with a PCR fragment. The fragment was obtained by PCR amplification using degenerate primers to randomize positions 366, 368 and 407 using standard techniques.

After 2 to 5 rounds of selection, the fraction of full length clones was 90%, 60%, 50% and 10%, respectively, as judged by agarose gel electrophoresis of single-stranded DNA. Phagemids containing full length clones were gel-purified after 5 rounds of selection. Two thousand transformants were obtained after retransforming XL1-BLUE™ cells (Stratagene).

A mean of > 10⁶ copies of each clone was used per round of panning. Thus, numerous copies of each clone in the library were likely available for selection, even though some deletion mutants arose during panning.

After 7 rounds of panning, the C_H3 mutants obtained approached a consensus amino acid sequence at the randomized residues. Virtually all clones had serine or threonine at residue 366 indicating a very strong preference for a β-hydroxyl at this position. A strong preference for hydrophobic residues was observed for residues 368 and 407, with valine and alanine predominating. Six different amino acid combinations were recovered at least twice, including the triple mutant, T366S:L368A:Y407V, which was recovered 11 times. None of these phage selectants has an identical sequence to a previously designed heterodimer, T366 W/Y407'A (Ridgway, J. B. B., *et al.*, (1996), *supra*. The phage selectants may be less tightly packed than the wild-type C_H3 homodimer as judged by a 40-80 ų reduction in total side chain volume of the domain interface residues.

C_H3 variants encoded on the expression plasmid pAK19 (Carter *et al.* 1992) were introduced into *E.coli* strain 33B6, expressed, and secreted from *E. coli* grown to high cell density in a fermentor. The T366S:L368A:Y407V mutant purified by DEAE-Sepharose FF, ABx and Resource S chromatography gave a single major band following SDS-PAGE. Other C_H3 variants were recovered with similar purity. The molecular masses of wild-type C_H3 and

T366S:L368A:Y407V, T366W and Y407A variants determined by high resolution electrospray mass spectrometry were as expected.

B. <u>Phage-selected heterodimer stability</u>. The stability of C_H3 heterodimers was first assessed by titrating corresponding phage with guanidine hydrochloride, followed by dilution and quantification of residual heterodimer by enzyme-linked immunosorbent assay (ELISA). The guanidine hydrochloride denaturation assay with C_H3-phage provides a means to screen selectants rapidly.

Phage were prepared from individual clones following 7 rounds of selection and also from the control vector, pRA1. Briefly, phagemids in XL1-BLUETM were used to inoculate 25 ml LB broth containing 50 μ g/ml carbenicillin and 10 μ g/ml tetracycline in the presence of 10⁹ pfu/ml M13K07 and incubated overnight at 37°C. The cells were pelleted by centrifugation (6000 g, 10 min, 4 °C). Phage were recovered from the supematantby precipitation with 5 ml 20 % (w/v) PEG, 2.5 M NaCl followed by centrifugation (12000 g, 10 min, 4 °C) and then resuspended in 1 ml PBS. 180 μ l 0-6 M guanidine hydrochloride in PBS was added to 20 μ l phage preparations and incubated for 5.0 min at approximately~25°C. Aliquots (20 μ l) of each phage sample were then diluted 10-fold with water. The presence of C_H3 heterodimerwas assayed by ELISA using 5B6-coated plates and detecting the phage with an anti-M13 polyclonal Ab conjugated to horseradish peroxidase, using o-phenylenediamine as the substrate. The reaction was quenched by the addition of 50 μ l 2.5 M H₂SO₄ and the absorbance measured at 492 nm. The absorbance data were plotted against the guanidine hydrochloride concentration during the melt and fitted to a 4 parameter model by a non-linear least squares method using Kaleidagraph 3.0.5 (Synergy Software).

The most frequently recovered heterodimer, T366W/T366'S:L368'A:Y407'V, is similar in stability to other phage-selected heterodimers. This phage-selected heterodimer is significantly more stable than the designed heterodimer, T366W/Y407'A but less stable than the wild-type C_H3 . All C_H3 variants, both individually and in combination, were found to be dimers by size exclusion chromatography under the conditions that these same molecules were studied by calorimetry (1.75 mg/ml, in phosphate-buffered saline (PBS)). The only exception was the T366S:L368A:Y407V mutant alone which had a slightly shorter retention time than C_H3 dimers.

A 1:1 mixture of T366W, protuberance, and T366S:L368A:Y407V, cavity, mutants melts with a single transition at 69.4 °C, consistent with subunit exchange and formation of a stable heterodimer. In contrast, the T366W protuberance homodimer is much less stable than the T366W/T366'S:L368'A:Y407'V protuberance-into-cavity heterodimer (ΔT_m =-15.0 °C). The T366S:L368A:Y407V cavity mutant on its own is prone to aggregate upon heating and does not undergo a smooth melting transition.

The designed cavity mutant, Y407A, melts at 58.8 °C and 65.4 °C in the absence and presence of the T366W protuberance mutant, respectively. This is consistent with subunit exchange and formation of a T366W/Y407'A heterodimerthat has greater stability than either T366W ($\Delta T_m = 11.0$ °C) or Y407A ($\Delta T_m = 6.6$ °C) homodimers. The phage-selectedheterodimer, T366W/T366'S:L368'A:Y407'V, is more stable than the designed heterodimer, T366W/Y407'A, ($\Delta T_m = 4.0$ °C), but is less stable than the wild-type C_H 3 homodimer ($\Delta T_m = -11.0$ °C).

C. Multimerization of a phage-selected antibody immunoadhesin (Ab/la) in vivo. Phage-selected and designed C_H3 mutants were compared in their ability to direct the formation of an Ab/la hybrid, anti-CD3/CD4-lgG *in vivo* (Chamow *et al.*, (1994), *supra*. This was accomplished by coexpression of humanized anti-CD3 light (L) and heavy chains together with CD4-lgG. Formation of heterodimers and homodimers was assessed by protein A purification followed by SDS-PAGE and scanning laser densitometry (Ridgway, *et al.*, (1996), *supra*). Comparable yields of Ab/la hybrid were recovered from cotransfections in which the anti-CD3 heavy chain contained the designed protuberance mutation, T366W, and the la contained either the phage-selected mutations, T366S:L368A:Y407V, or designed cavity mutation, Y407A (Fig. 3).

Phage-selected and designed C_H3 mutants were next evaluated in their propensity to form homodimers. The protuberance mutation, T366W, is apparently very disruptive to homodimerization since cotransfection of corresponding antibody heavy and light chains leads to an excess of HL monomers (may include non disulfide-bonded IgG) over IgG. In contrast, IgG but no HL monomers are observed for the same antibody containing wild-type C_H3 domains. The cavity mutations, T366S:L368A:Y407V, are somewhat disruptive to homodimerization since transfection of the corresponding phagemid leads to a mixture of predominantly la dimers with some la monomers. The cavity mutation, Y407A, is minimally disruptive to homodimerization as judged by the presence of la dimers but no la monomers following transfection of the corresponding phagemid.

The phage display selection strategy described herein allows the selection in favor of C_H3 mutants that form stable heterodimers and selection against mutants that form stable homodimers. The counter selection against homodimers occurs because "free" C_H3 mutants will compete with the flagged C_H3 knob mutant for binding to available C_H3 mutant-gene III fusion protein. The free C_H3 mutants arise as a result of the amber mutation between the natural C_H3 gene and M 13 gene III. In an amber suppressor host such as XLI-Blue, both C_H3 -gene III fusion protein and corresponding free C_H3 will be secreted.

Guanidine hydrochloridedenaturation proved to be a useful tool for the preliminary screening of the stability of C_H3 heterodimers on phage. Phage maintain infectivity for E, coli even after exposure to 5 M guanidine hydro-

32

10

5

20

15

25

30

35

40

45

50

55

chloride (Figini *et al.*, J. Mol. Biol. <u>239</u>:68-78 (1994)). Thus, guanidine may also be useful to increase the stringency of mutant selection.

Rational design and screening of phage display libraries are complementaryapproachesto remodeling a domain interface of a homodimer to promote heterodimerization. In the case of C_H3 domains, designed mutants identified domain interface residues that could be recruited to promote heterodimerization. Phage display was then used here to search permutations of 3 residues neighboring a fixed protuberance for combinationsthat most efficiently form heterodimers. Phage selectants are useful to facilitate further rational redesign of the domain interface, while the phage selection strategy described herein demonstrates its usefulness for remodeling protein-protein interfaces.

5

10

15

20

25

30

40

45

50

55

Example 4: Generation and assembly of heteromultimeric antibodies or antibody/immunoadhesins having common light chains

[0172] The following example demonstrates preparation of a heteromultimeric bispecific antibody sharing the same light chain according to the invention and the ability of that antibody to bind its target antigens.

A. Identification of antibodies that share the same light chain: Comparison of antibody libraries raised to eleven antigens.

[0173] A large human single chain Fv (scFv) antibody library (Vaughan $et\,al.$ (1996), supra) was panned for antibodies specific for eleven antigens including Axl(human receptor tyrosine kinase ECD), GCSF-R (human granulocyte colony stimulating factor receptor ECD), IgE (murine IgE), IgE-R (human IgE receptor α -chain), MPL (human thrombopoietin receptor tyrosine kinase ECD), MusK (human muscle specific receptor tyrosine kinase ECD), NpoR (human orphan receptor NpoR ECD), Rse (human receptor tyrosine kinase, Rse, ECD), HER3 (human receptor tyrosine kinase HER3/c-erbB3 ECD), Ob-R (human leptin receptor ECD), and VEGF (human vascular endothelial growth factor) where ECD refers to the extracellular domain. The nucleotide sequence data for scFv fragments from populations of antibodies raised to each antigen was translated to derive corresponding protein sequences. The V_L sequences were then compared using the program "align" with the algorithm of Feng and Doolittle (1985, 1987, 1990) to calculate the percentage identity between all pairwise combinations of chains (Feng, D.F. and Doolittle, R.F. (1985) J. Mol. Evol. $\underline{21}$:112-123; Feng, D.F. and Doolittle, R.F. (1987) J. Mol. Evol. $\underline{25}$:351-360; and Feng, D.F. and Doolittle, R.F. (1990) Methods Enzymol. $\underline{183}$:375-387). The percent sequence identity results of each pairwise light chain amino acid sequence comparison were arranged in matrix format (see Appendix).

[0174] For most pairwise comparisons, at least one common light chain sequence was found. Table 5 is a comparison of the V_L chains showing the frequencies of scFv sharing identical light chains (100% identity) determined by alignment of 117 V_L amino acid sequences. For example, the entry 4/9 (HER3 x Ob-R, highlighted in a black box), denotes that 4 clones that bind HER3 were found to share their V_L sequence with one or more anti-Ob-R clones, whereas 9 clones binding the Ob-R share their V_L sequence with one or more anti-HER3 clones. The entries on the diagonal represent the number of antibody clones within a population that share a V_L sequence with one or more clones in the population. For example, examination of the MPL clones revealed 5 clones that shared their V_L sequence with one or more other MPL clones. In the cases where no common light chain sequence was observed, such as for (IgE x AxI) or (NpoR x IgE-R), the number of fragments compared for at least one specificity was very small (5 or less). Given the number of common light chains found, it is likely that common light chains can be found for any V_L comparison if a sufficient number of clones are compared.

[0175] The amino acid sequences of light chains were examined for the positions of amino acid residue differences when the sequence identity relative to a chosen common light chain was 98% and 99%. Fig. 4 is a comparison of V_L sequences of eight different antibodies with specificities for AxI (clone Ax1.78), Rse (clones Rse.23, Rse.04, Rse.20, and Rse.15), IgER (clone IgER.MAT2C1G11), Ob-R (clone obr.4), and VEGF (clone vegf.5). The position of the antigen binding CDR residues according to a sequence definition (Kabat, G. A., *et al.* (1991) *supra*) or structural definition (Chothia and Lesk, (1987) J. Mol. Biol. 196:901-917) are shown by underlining and #, respectively. Light chain residues that differ from the Ax1.78 sequence are shown by double underlining. Of the 9 light chains compared, 6 are identical. The light chains of Rse.04 and obr.4 (approximately 99% sequence identity) differ by one residue outside of the antigen binding CDRs. The light chain of Rse.20 (approximately98% sequence identity) differs by two residues outside of the antigen binding CDRs. The amino acid residue changes may have little or no affect on antigen binding. Thus, the sequence similarity of these light chains makes them candidates for the common light chain of the invention. Alternatively, according to the invention, such light chains having 98-99% sequence identity with the light chain of a prospective paired scFv (Ax1.78, for example) may be substituted with the paired light chain and retain binding specificity.

B. Identification of antibodies that share the same light chain and construction of a bispecific antibody sharing that light chain: Anti-Ob-R/Anti-HER3.

[0176] ScFv fragments that bound human leptin receptor (Ob-R) or the extracellular domain of the HER3/c-erbB3 gene product (HER3) were obtained by three rounds of panning using a large human scFv phage library (Vaughan *et al.* (1996), *supra*). Leptin receptor-IgG and HER3-IgG (10 μg in 1 ml PBS were used to coat separate Immunotubes (Nunc; Maxisorp) overnight at 4°C. Panning and phage rescue were then performed as described by Vaughan *et al.* (1996), *supra*, with the following modifications. A humanized antibody, huMAb4D5-8 (Carter, P. *et al.* (1992) PNAS USA 89:4285-4289) or humanized anti-IgE (Presta, L. *et al.* (1993) J. Immunol. 151:2623-2632)at a concentration of 1 mg/ml was included in each panning step to absorb Fc-binding phage. In addition, panning in solution (Hawkins, R. E., *et al.* (1992) J. Mol. Biol. 226:889-896) was also used to identify scFv binding leptin receptor. The leptin receptor was separated from the Fc by site-specific proteolysis of leptin receptor-IgG with the engineered protease, Genenase (Carter, P., *et al.* (1989) Proteins: Structure, Function and Genetics 6:240-248) followed by protein A Sepharose chromatography. The leptin receptorwas biotinylated and used at a concentration of 100 nM, 25 nM and 5nM for the first, second, and third rounds of panning, respectively. Phage binding biotinylated antigen were captured using streptavidincoated paramagnetic beads (Dynabeads, Dynal, Oslo, Norway).

[0177] Clones from rounds 2 and 3 of each panning were screened by phage and scFv ELISA using the corresponding antigen and also a control immunoadhesinor antibody. The diversity of antigen-positive clones was analyzed by PCR-amplification of the scFv insert using the primers, fdtetseq and PUC reverse (Vaughan et al. (1996), supra) and by digestion with BstNI (Marks et al. (1991) supra). One to five clones per BstN1 fingerprint were then cycle-sequencedusing fluorescent dideoxy chain terminators (Applied Biosystems) using PCR heavy link and myc seq 10 primers (Vaughan et al. (1996), supra). Samples were analyzed using an Applied Biosystems Automated DNA Sequencer and sequences analyzed using SeqEd. It is also noted that the quanidine hydrochloride antibody denaturation and in vitro chain shuffling method of Figini combined with phage display selection is useful as a method of selecting antibodies having the same light chain (Figini, M. et al. (1994), supra).

[0178] Using the method described above, eleven different anti-HER3 clones and 18 anti-Ob-R clones (1I form panning using coated antigen and 7 from panning with biotinylated antigen) were obtained. The clones were sequenced by standard techniques to determine the sequences of the light chains associated with each binding domain (Fig. 5). The sequences are the V_H and common V_L sequences of the anti-Ob-R clone 26 and anti-HER3 clone 18 used to construct a bispecific antibody (see below). The residues are numbered according to (Kabat, E. A., *et al.* (1991) *supra*). The position of the antigen binding CDR residues according to a sequence definition (Kabat et al. (1991) *supra*) or structural definition (Chothia and Lesk, (1987) J. Mol. Biol. (1987) 196:901-917) are shown by underlining and overlining, respectively. Identity between residues in the V_H sequences is indicated by *.

[0179] The sequences of the light chains were compared for multiple anti-HER3 clones relative to multiple anti-Ob-R clones (Fig. 8 and Table 5). It was observed that four out of eleven anti-HER3 clones share identical V_L with one or more anti-Ob-R receptorclones. Conversely, nine out of eighteen anti-Ob-R clones share the same V_L as one of the anti-HER3 clones (See Table 5, blackened box).

	•
Table 5	Shared V _L usage by scFv against different target antigens

Antigen	*	Ax	SCS	1gE	lgE-R	MPL	MusK	MusK NpoR	Rse	HER3	Ob-R	VEGF
Specificity	ScF		F-R									
	>											
Axi	12	2	2/2	0/0	1/1	2/3	1/1	0/0	3/5	272	2/5	1/1
GCSF-R			0	1/1	272	2/3	1/1	2/2	2/3	2/2	3/3	2/3
1gE	7			0		1/1	0/0	1/1	1/1	1/1	1/1	0/0
lgE-R	4				0	1/1	0/0	1/1	2/3	1/1	1/1	1/1
MPL	23					\$	5/3	3/2	8/9	7/5	6/5	2/2
MusK	m						0	1/1	1/2	2/2	1/1	1/2
NpoR .	٧٠							0	1/1	2/2	2/2	1/2
Rse	20								7	7/4	2/8	2/1
HER3			٠							m	419	4/4
Ob-R	8		•								۲.	172
VEGF	∞			•								7

[0180] Construction of anti-Ob-R/anti-HER3, a bispecific antibody having a common light chain was performed as follows. Altered C_H3 first and second polypeptides having the complementary protuberances and cavities as well as the non-naturally occurring disulfide bonds between the first and second polypeptides were used in the construction of a Fc-containing bispecific antibody. The V_L from anti-Ob-R clone #26 and anti-HER3 clone #18, which clones share the same light chain, as well as the heavy chains from each antibody were used to prepare the bispecific antibody according to the procedures disclosed herein.

[0181] This antibody had an electrophoretic mobility shift in apparent molecular weight relative to a bispecific antibody that differed only by a lack of alterations for generating non-natural disulfide bonds. An 8% SDS-PAGE gel of heterodimeric antibody variants with and without non-naturally occurring disulfide bonds showed a mobility shift from approximately 230 apparent MW for wild type heterodimer to approximately 200 apparent MW for a heterodimerhaving one non-natural disul fide bond. The MW shift was sufficient to allow determination of the percent of each variant that successfully formed the non-natural disulfide bond.

[0182] The binding specificity for both Ob-R and for HER3 of the bispecific antibody is tested by standard ELISA procedures such as the following method. Ob-R binding is demonstrated in an ELISA assay with Ob-R present as an

Ob-R-Ig fusion protein. The Ob-R-Ig fusion protein is coated onto the well of a 96-well microtitre plate and the bispecific antibody is added.

[0183] The well is washed several times to eliminate non-specific binding to Ob-R-Ig. As a second component in the same assay, a biotinylated HER3-Ig fusion protein is added and detected by means of streptavidin-horseradishperoxidasecomplex binding to the biotinylated HER3-Ig fusion protein. Binding is detected by generation of a color change upon addition of hydrogen peroxide and TMB peroxidase substrate (Kirkegaard and Perry Laboratories, Gaithersburg, MD).

5

10

15

20

25

30

40

45

50

55

[0184] Under the conditions just described, the binding of a bispecific antibody to both Ob-R-Ig and to HER3-Ig would be observed as detectable label immobilized on the surface of the microtitre well due to the formation of a complex comprising immobilized Ob-R-Ig/bispecific antibody/HER3-Ig biotin/detectably labeled streptavidin. Antibodies that bind Ob-R-Ig, but not HER3-Ig, do not form the above complex, providing a negative result. Similarly, antibodies that bind to HER3-Ig, but not Ob-R-Ig, do not form the above complex and provide a negative result. In contrast, the bispecific antibody expected to bind both Ob-RIg and HER3-Ig, forms the complex yielding a positive result in the assay, demonstrating that the bispecific antibody, having a common light chain, binds both HER3 and Ob-R.

[0185] Expression and purification of the anti-(Ob-R/HER3) bispecific antibody was performed as follows. Human embryonic kidney 293S cells were transfected with three plasmid DNAs each separately encoding anti-Ob-R heavy chain, anti-HER3 heavy chain, or the light chain from clone 26 or 18 that was common to each of the antibodies, as described *supra*. For each transfection, the ratio of heavy chain-encoding DNA to light chain-encoding DNA was 1:3 so that light chain would not be limiting for assembly of anti-Ob-R/anti-HER3 bispecific antibody. Both heavy chains were transfected in a 1:1 ratio with respect to each other. 12 μg of total plasmid DNA was then co-transfected into 293S cells by means of calcium phosphate precipitation (Gorman, C., DNA Cloning, Vol. H, D.M. Glover, ed., IRL Press, Oxford, p. 143 (1985)). The cells were washed with PBS prior to adding growth media intended to enhance protein expression. Fc-containing proteins were purified from cell supernatants using immobilized protein A (ProSep A, BioProcessing Ltd., UK) and buffer-exchanged into PBS. Iodoacetamidewas added to protein preparations to a final concentration of 50 mM to prevent reshuffling of disulfide bonds.

[0186] As an additional example, expression and purification of an anti-(CD3/CD4) antibody/immunoadhesin was performed as follows. Human embryonic kidney 293S cells were transfected with three plasmid DNAs, each plasmid separately encoding anti-CD3 light chain, anti-CD3 lgG₁ heavy chain, or anti-CD4 lgG₁ immunoadhesin. For each transfection, the ratio of light chain-encoding DNA to heavy chain-encoding DNA was 3:1 so that light-chain would not be limiting for assembly of anti-CD3 lgG. Additionally, because the immunoadhesin is poorly expressed, the ratio of immunoadhesin encoding plasmid was added in excess to heavy chain encoding plasmid. The ratios tested ranged from 3:1:3 through 8:1:3 for immunoadhesin:heavychain:light chain phagemids. 10 µg total plasmid DNA were then co-transfected into 293S cells by means of calcium phosphate precipitation (Gorman, C. (1985), *supra*), washing cells with PBS prior to transfection. Fc-containing proteins were purified from cell supernatants using immobilized protein A (ProSep A, BioProcessingLtd., UK) and buffer-exchanged into PBS. Iodoacetamide was added to protein preparations to a final concentration of 50 mM to prevent reshuffling of disulfide bonds.

[0187] In each of the above preparations, protein samples were electrophoresed on 8% polyacrylamide gels (Novex) and visualized by staining with Serva blue. Gels were de-stained leaving a faint background in an effort to allow visualization and quantitation of minor contaminants. Dried gels were scanned with the scanning densitometer (GS-670, BioRad) and protein products were quantitated with Molecular Analyst software.

[0188] Non-natural(engineered)disulfide bonds introduced into the C_H3 domain has been disclosed herein to enhance heterodimerformation. One pair of polypeptides, K392C/D399'C, enhanced heterodimer formation by generating up to 76% heterodimer (Table 4, variant v6). Moreover, when the presence of an inter-chain disulfide bond was combined with the protuberance-into-cavity technology, approximately 95% heterodimer was obtained (Table 4 variants v11, v12, and v16). Thus, the method of the invention of increasing specific protein/protein interaction between the first and second polypeptides of a bispecific antibody increases the yield of desired heteromultimer and minimizes the formation of undesired heteromultimersor homomultimers.

[0189] In addition, the method of characterizing the product heteromultimers by electrophoretic mobility analysis allows for the determination of the relative amount of desired heteromultimers relative to undesired products.

[0190] Selection of a common light chain as described herein further increases yield of the desired heteromultimer by eliminating the possibility of mispairing between variable heavy chains and light chains of a multispecific antibody.

C. Identification of antibodies that share the same light chain and construction of a bispecific antibody sharing that light chain: Anti-Mpl/Anti-HER3.

[0191] Identification, construction and expression of another bispecific antibody of the invention is demonstrated herein. The methods described in Parts A and B of this example were utilized for the preparation of the anti-Mpl/anti-HER3 bispecific antibody.

[0192] Using the methods described in Section A of this example (Comparison of antibody libraries raised to eleven antigens), *supra*, the V_H and V_L amino acid sequences of the anti-HER3 scFv were compared with 23 scFv that bind to the human thrombopoietin receptor, c-Mpl. Five of the eleven anti-HER3 clones share an identical V_L amino acid sequence with one or more Mpl-binding clones. Conversely, seven out of twenty-three anti-Mpl scFv shared the same V_L as one of the anti-HER3 clones (see Table 5, *supra*, open box). In contrast, the V_H amino acid sequences were much more diverse, with an identity level of 40 to 90% between any anti-Mpl and anti-HER3 clone.

[0193] The anti-Mpl scFv, 12B5 (Genbank accession number AF048775; SEQ ID NO:27) and anti HER3 scFv clone H6 (Genbank accession number AF048774; SEQ ID NO:28) utilize identical V_L sequences and substantially different V_H sequences. These scFv fragments were used to construct the anti-Mpl/anti-HER3 bispecific IgG antibody capable of efficient heterodimerizationdue to the shared light chain as well as through the use of knobs-into-holes mutations (described herein) and an engineered disulfide bond between the C_H 3 domains. Antibodies that share the same L chain were chosen to circumvent the problem of L chains pairing with non-cognate H chains. Two naturally occurring hinge region disulfide bonds were also present. The common L chain was cotransfected with the two H chains containing the C_H 3 mutations from variant v11. The IgG products were purified by protein A affinity chromatography and analyzed by SDS-PAGE using standard techniques.

[0194] The bispecific IgG antibody (BsIgG) preparation gave rise to a single major band showing greater mobility than IgG containing wild-type C_H3 domains. This increase in electrophoreticmobilitywas consistent with the formation of the engineered disulfide bond in the BsIgG forming a more compact protein species.

[0195] The ability of the engineered anti-Mpl/anti-HER3 BslgG antibody to bind both Mp1 and HER3 ECD antigens was assessed using an ELISA as follows. Using PBS buffer in all steps, individual wells of a 96 well plate (Maxisorp, Nunc) were coated overnight with HER3-lgG or Mp1-lgG at 5 μg/ml, washed and then blocked for 1 hour with 0.5% (w/v) BSA. The primary antibodies were the anti-Mp1 x anti-HER3 BslgG containing the mutations, Y349C:T366S: L368A:Y407V/T366'W:S354'C and corresponding parental anti-Mp1 oranti-HER3 lgG with mutated Fc regions. The primary antibodies (1 μg/mL) were individually incubated at 2 h at 23° C with biotinylated HER3-lgG and a 1:5000 dilution of streptavidin-horse radish peroxidase conjugate (Boehringer Mannheim) and then added to the wells and incubated for an additional 1 h at 23° C. Peroxidase activity was detected with TMB reagents as directed by the vendor (Kirkegaard and Perry Laboratories, Inc., Gaithersburg, MD).

[0196] As anticipated, the anti-Mp1/anti-HER3Bs1gG bound efficiently and simultaneously to each Mpl and HER3 ECD antigens individually as well as to both antigens simultaneously. By contrast, the parental anti-Mpi and parental anti-HER3 IgG bound only to their corresponding cognate antigen (Fig. 6).

D. Antibodies containing an engineered Fc region are capable of efficient antibody-dependent cell-mediated cytotoxicity.

[0197] To demonstrate that the engineered F_c region (C_H 3 mutations, supra) utilized in generating the exemplified bispecific antibodies of the invention is capable of efficient antibody-dependent cell-mediated cytotoxicity (ADCC), the following experiment was performed.

[0198] The C_H3 mutations maintain the ability to support efficient antibody-dependent cell-mediated cytotoxicity (AD-CC) as demonstrated using the method of Lewis, G.D. et al. (Lewis, G.D, et al. (1993) Cancer Immunol. Immunother. 37:255-263). Briefly, cytotoxicity assays were performed with ⁵¹Cr-labeled SK-BR-3 and HBL-100 target cells (ATCC accession numbers HTB-30 and 45509, respectively)and human peripheral blood lymphocytes as effectorcells. However, unlike Lewis et al., the lymphocytes were not activated with IL-2.

[0199] The C_H3 mutations S354:T366W and Y349:T366S:L368A:Y407V were introduced separately into the H chain of the humanized anti-HER2 antibody, huMAb4D5-5 prepared by Carter et al. (Carter, P. et al. (1992) PNAS USA <u>89</u>: 4285-4289). Antibodies containing remodeled and wild-type Fc regions had similar potency in ADCC with the HER2-overexpressingbreast cancer cell line, SK-BR-3 (Fig. 7). Both remodeled and wild-type antibodies showed comparable, low activity against the normal breast epithelial cell line. The effects in the H-chain are independent of the binding domains, predicting that these BsIgG's will function in antibody-dependent cell-mediated cytotoxicity.

50 SEQUENCE LISTING

[0200]

5

15

20

25

30

45

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: GENENTECH, INC.
 - (ii) TITLE OF INVENTION: A METHOD FOR MAKING MULTISPECIFIC ANTIBODIES HAVING HETEROM-

ULTIMERIC AND COMMON COMPONENTS

	(iii) NUMBER OF SEQUENCES: 28
5	(iv) CORRESPONDENCE ADDRESS:
10	 (A) ADDRESSEE: Genentech, Inc. (B) STREET: 1 DNA Way (C) CITY: South San Francisco (D) STATE: California (E) COUNTRY: USA (F) ZIP: 94080
4.5	(v) COMPUTER READABLE FORM:
15 20	(A) MEDIUM TYPE: 3.5 inch, 1.44 Mb floppy disk(B) COMPUTER: IBM PC compatible(C) OPERATING SYSTEM: PC-DOS/MS-DOS(D) SOFTWARE: WinPatin (Genentech)
	(vi) CURRENT APPLICATION DATA:
25	(A) APPLICATION NUMBER: PCT/US98/08762(B) FILING DATE: 30-Apr-1998(C) CLASSIFICATION:
	(vii) PRIOR APPLICATION DATA:
30	(A) APPLICATION NUMBER: 08/850058 (B) FILING DATE: 02-MAY-1997
	(viii) ATTORNEY/AGENT INFORMATION:
35	(A) NAME: Conley, Deirdre L. (B) REGISTRATION NUMBER: 36,487 (C) REFERENCE/DOCKET NUMBER: P1099R2PCT
	(ix) TELECOMMUNICATION INFORMATION:
40	(A) TELEPHONE: 650/225-2066 (B) TELEFAX: 650/952-9881
	(2) INFORMATION FOR SEQ ID NO:1:
45	(i) SEQUENCE CHARACTERISTICS:
50	(A) LENGTH: 36 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single,(D) TOPOLOGY: Linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
	(,
55	CTCTTCCCGA GATGGGGGCA GGGTGCACAC CTGTGG 36

(2) INFORMATION FOR SEQ ID NO:2:

	(i) SEQUENCE CHARACTERISTICS:
5	(A) LENGTH: 21 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
10	CTCTTCCCGA CATGGGGGCA G 21
	(2) INFORMATION FOR SEQ ID NO:3:
15	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 21 base pairs
20	(B) TYPE: Nucleic Acid
	(C) STRANDEDNESS: Single
	(D) TOPOLOGY: Linear
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
	GGTCATCTCA CACCGGGATG G 21
30	(2) INFORMATION FOR SEQ ID NO:4:
	(i) SEQUENCE CHARACTERISTICS:
35	(A) LENGTH: 24 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
	CTTGGTCATA CATTCACGGG ATGG 24
45	(2) INFORMATION FOR SEQ ID NO:5:
40	(i) SEQUENCE CHARACTERISTICS:
50	(A) LENGTH: 30 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:
55	CTCTTCCCGA GATGGGGGAC AGGTGTACAC 30
	(2) INFORMATION FOR SEQ ID NO:6:

	(i) SEQUENCE CHARACTERISTICS:
5	(A) LENGTH: 21 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:
10	GCCGTCGGAA CACAGCACGG G 21
15	(2) INFORMATION FOR SEQ ID NO:7:
20	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:
25	CTGGGAGTCT AGAACGGGAG GCGTGGTACA GTAGTTGTT 39
	(2) INFORMATION FOR SEQ ID NO:8:
30	(i) SEQUENCE CHARACTERISTICS:
<i>35</i>	(A) LENGTH: 33 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
40	GTCGGAGTCT AGAACGGGAG GACAGGTCTT GTA 33
	(2) INFORMATION FOR SEQ ID NO:9:
45	(i) SEQUENCE CHARACTERISTICS:
50	(A) LENGTH: 21 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:
55	GTCGGAGTCT AGACAGGGAG G 21
55	(2) INFORMATION FOR SEQ ID NO:10:
	(i) SEQUENCE CHARACTERISTICS:

5	(A) LENGTH: 21 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
10	GCCGTCGGAG CTCAGCACGG G 21
	(2) INFORMATION FOR SEQ ID NO:11:
15	(i) SEQUENCE CHARACTERISTICS:
20	(A) LENGTH: 24 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:
	GGGAGGCGTG GTGCTGTAGT TGTT 24
25	(2) INFORMATION FOR SEQ ID NO:12:
	(i) SEQUENCE CHARACTERISTICS:
30	(A) LENGTH: 38 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:
	GTTCAGGTGC TGGGCTCGGT GGGCTTGTGT GAGTTTTG 38
40	(2) INFORMATION FOR SEQ ID N0:13:
	(i) SEQUENCE CHARACTERISTICS:
45	(A) LENGTH: 821 base pairs(B) TYPE: Nucleic Acid(C) STRANDEDNESS: Single(D) TOPOLOGY: Linear
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:
55	

TCTGGCTGCA CACTACGGCC AGCCGCGGA ACCTCAGGTG TATACCCTGC 1 CACCGTCTCG AGAAGAAACC AGGTCTCTC GTGGTGCCTG 1 GTCAAAGGTT TCTATCCGAG CGATATCGCC GTGGAATGGG AAAGCAACGG 2 GCTCCTTCTT TCTGTATTCG AAACCACTCC ACCGGTGCTG GATTCTGATG 2 GCTCCTTCTT TCTGTATTCG AAGCTGACCG TTGACAAAAG CCGTTGGCAG 3 CAAGGCAACG TTTCAGCTG TCTGTTATG CACGAGGCCT TGCACAACCA 3 CTACACCCAG AAAAGCCTGT CCCTGTCTCC CGGGAAATAA GCTGAGGCTC 4 CTCTAGAGGT TGAGGTGATT TTATGAAAAA GAATATCGCA TTTCTTCTTG 4 CAACCACGA AGGTGTACCA CCTGCCCCCA TCCCGGGAAG AGATGACCAA 5 CGAGAACCAC AGGTGTACCA CCTGCCCCCA TCCCGGGAAG AGATGACCAA 5 CGACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTT CCCAGCGACA 6 TCCCCGTGGA GTGGGAGACC CGGAGAACAA CTACAAGACC 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7 TGATGCATGA GGCTTCTGCAC AACCACTACA CGCAGAAGAG CCTCTCCCTG 7		AACGCGTACG	CTCTGAAAAT	GGCGGACCCG	AACCGTTTTC	GTGGTAAAGA	50
GTCAAAGGTT TCTATCCGAG CGATATCGCC GTGGAATGGG AAAGCAACGG 2 TCAACCGGAA AACAACTACA AAACCACTCC ACCGGTGCTG GATTCTGATG 2 GCTCCTTCTT TCTGTATTCG AAGCTGACCG TTGACAAAAG CCGTTGGCAG 3 CAAGGCAACG TTTTCAGCTG TTCTGTTATG CACGAGGCCT TGCACAACCA 3 CTACACCCAG AAAAGCCTGT CCCTGTCTCC CGGGAAATAA GCTGAGGCTC 4 CTCTAGAGGT TGAGGTGATT TTATGAAAAA GAATATCGCA TTTCTTCTTG 4 CATCTATGTT CGTTTTTCT ATTGCTACAA ACGCGTACGC TGGGCAGCCC 5 CGAGAACCAC AGGTGTACAC CCTGCCCCCA TCCCGGGAAG AGATGACCAA 5 GAACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTAT CCCCAGCGACA 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTCT 7 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTCT 7 ACACCTCCCC TGCTGGACTC CGACGGCTCC TTCTTCCTCT TCATGCTCCC 7	5	TCTGGCTGCA	CACTACGGCC	AGCCGCGGGA	ACCTCAGGTG	TATACCCTGC	100
TCAACCGGAA AACAACTACA AAACCACTCC ACCGGTGCTG GATTCTGATG 2 GCTCCTTCTT TCTGTATTCG AAGCTGACCG TTGACAAAAG CCGTTGGCAG 3 CAAGGCAACG TTTTCAGCTG TTCTGTTATG CACGAGGCCT TGCACAACCA 3 CTACACCCAG AAAAGCCTGT CCCTGTCTCC CGGGAAATAA GCTGAGGCTC 4 CTCTAGAGGT TGAGGTGATT TTATGAAAAA GAATATCGCA TTTCTTCTTG 4 CATCTATGTT CGTTTTTCT ATTGCTACAA ACGCGTACGC TGGGCAGCCC 5 CGAGAACCAC AGGTGTACAC CCTGCCCCA TCCCGGGAAG AGATGACCAA 5 CACCGTGGA GTGGGAGGC CCTGGTCAA AGGCTTCTAT CCCAGCGACA 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7		CACCGTCTCG	AGAAGAAATG	ACTAAAAACC	AGGTCTCTCT	GTGGTGCCTG	150
GCTCCTTCTT TCTGTATTCG AAGCTGACCG TTGACAAAAG CCGTTGGCAG 3 CAAGGCAACG TTTTCAGCTG TTCTGTTATG CACGAGGCCT TGCACAACCA 3 CTACACCCAG AAAAGCCTGT CCCTGTCTCC CGGGAAATAA GCTGAGGCTC 4 CTCTAGAGGT TGAGGTGATT TTATGAAAAA GAATATCGCA TTTCTTCTTG 4 CATCTATGTT CGTTTTTCT ATTGCTACAA ACGCGTACGC TGGGCAGCCC 5 CGAGAACCAC AGGTGTACAC CCTGCCCCA TCCCGGGAAG AGATGACCAA 5 GAACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTAT CCCAGCGACA 6 TCGCCGTGGA GTGGGAGAGC AATGGGCAGC CGGAGAACAA CTACAAGACC 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7		GTCAAAGGTT	TCTATCCGAG	CGATATCGCC	GTGGAATGGG	AAAGCAACGG	200
CAAGGCAACG TTTTCAGCTG TTCTGTTATG CACGAGGCCT TGCACAACCA 3 CTACACCCAG AAAAGCCTGT CCCTGTCTCC CGGGAAATAA GCTGAGGCTC 4 CTCTAGAGGT TGAGGTGATT TTATGAAAAA GAATATCGCA TTTCTTCTTG 4 CATCTATGTT CGTTTTTCT ATTGCTACAA ACGCGTACGC TGGGCAGCCC 5 CGAGAACCAC AGGTGTACAC CCTGCCCCCA TCCCGGGAAG AGATGACCAA 5 GAACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTAT CCCAGCGACA 6 TCGCCGTGGA GTGGGAGAGC AATGGGCAGC CGGAGAACAA CTACAAGACC 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTCT 7 30 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7	10	TCAACCGGAA	AACAACTACA	AAACCACTCC	ACCGGTGCTG	GATTCTGATG	250
CTACACCCAG AAAAGCCTGT CCCTGTCTCC CGGGAAATAA GCTGAGGCTC 4 CTCTAGAGGT TGAGGTGATT TTATGAAAAA GAATATCGCA TTTCTTCTTG 4 CATCTATGTT CGTTTTTCT ATTGCTACAA ACGCGTACGC TGGGCAGCCC 5 CGAGAACCAC AGGTGTACAC CCTGCCCCCA TCCCGGGAAG AGATGACCAA 5 GAACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTAT CCCAGCGACA 6 TCGCCGTGGA GTGGGAGAGC AATGGGCAGC CGGAGAACAA CTACAAGACC 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7		GCTCCTTCTT	TCTGTATTCG	AAGCTGACCG	TTGACAAAAG	CCGTTGGCAG	300
CTCTAGAGGT TGAGGTGATT TTATGAAAAA GAATATCGCA TTTCTTCTTG 4 CATCTATGTT CGTTTTTCT ATTGCTACAA ACGCGTACGC TGGGCAGCCC S CGAGAACCAC AGGTGTACAC CCTGCCCCCA TCCCGGGAAG AGATGACCAA S GAACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTAT CCCAGCGACA G TCGCCGTGGA GTGGGAGAGC AATGGGCAGC CGGAGAACAA CTACAAGACC G ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7	15	CAAGGCAACG	TTTTCAGCTG	TTCTGTTATG	CACGAGGCCT	TGCACAACCA	350
CATCTATGTT CGTTTTTCT ATTGCTACAA ACGCGTACGC TGGGCAGCCC S CGAGAACCAC AGGTGTACAC CCTGCCCCCA TCCCGGGAAG AGATGACCAA S GAACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTAT CCCCAGCGACA 6 TCGCCGTGGA GTGGGAGAGC AATGGGCAGC CGGAGAACAA CTACAAGACC 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7		CTACACCCAG	AAAAGCCTGT	CCCTGTCTCC	CGGGAAATAA	GCTGAGGCTC	400
CATCTATGTT CGTTTTTCT ATTGCTACAA ACGCGTACGC TGGGCAGCCC S CGAGAACCAC AGGTGTACAC CCTGCCCCCA TCCCGGGAAG AGATGACCAA S GAACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTAT CCCAGCGACA 6 TCGCCGTGGA GTGGGAGAGC AATGGGCAGC CGGAGAACAA CTACAAGACC 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7		CTCTAGAGGT	TGAGGTGATT	TTATGAAAAA	GAATATCGCA	TTTCTTCTTG	450
GAACCAGGTA AGCTTGTACT GCCTGGTCAA AGGCTTCTAT CCCAGCGACA 6 TCGCCGTGGA GTGGGAGAGC AATGGGCAGC CGGAGAACAA CTACAAGACC 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7	20	CATCTATGTT	CGTTTTTTCT	ATTGCTACAA	ACGCGTACGC	TGGGCAGCCC	500
TCGCCGTGGA GTGGGAGAGC AATGGGCAGC CGGAGAACAA CTACAAGACC 6 ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7		CGAGAACCAC	AGGTGTACAC	CCTGCCCCCA	TCCCGGGAAG	AGATGACCAA	550
ACGCCTCCCG TGCTGGACTC CGACGGCTCC TTCTTCCTCT ACAGCTTTCT 7 CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7	25	GAACCAGGTA	AGCTTGTACT	GCCTGGTCAA	AGGCTTCTAT	CCCAGCGACA	600
CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7		TCGCCGTGGA	GTGGGAGAGC	AATGGGCAGC	CGGAGAACAA	CTACAAGACC	650
CACCGTCGAC AAGAGCAGGT GGCAGCAGGG GAACGTCTTC TCATGCTCCG 7		ACGCCTCCCG	TGCTGGACTC	CGACGGCTCC	TTCTTCCTCT	ACAGCTTTCT	700
TGATGCATGA GGCTCTGCAC AACCACTACA CGCAGAAGAG CCTCTCCCTG 8	30	CACCGTCGAC	AAGAGCAGGT	GGCAGCAGGG	GAACGTCTTC	TCATGCTCCG	750
		TGATGCATGA	GGCTCTGCAC	AACCACTACA	CGCAGAAGAG	CCTCTCCCTG	800

35

40

45

TCTCCGGGTA AATAGGGGCC C 821

(2) INFORMATION FOR SEQ ID NO:14:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 50 amino acids

(B) TYPE: Amino Acid

(D) TOPOLOGY: Linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

50

		Ser 1	Asn	Arg	Phe	Ser 5	GTA	Ser	Lys	Ser	Gly 10	Asn	Thr	Ala	Ser	Leu 15
5		Thr	Ile	Ser	Gly	Leu 20	Gln	Ala	Glu	Asp	Glu 25	Ala	Asp	Tyr	Tyr	Cys 30
10		Ser	Ser	Tyr	Thr	Thr 35	Arg	Ser	Thr	Arg	Val 40	Phe	Gly	Gly	Gly	Thr 45
		Lys	Leu	Thr	Val	Leu 50										
15	(2) INFORMA	TION	FOR S	SEQ II	O NO:	15:										
	(i) SEQU	ENCE	CHA	RACTI	ERIST	ICS:										
20	(B) T	ENGT YPE: /	Amino	Acid												
	(xi) SEQU	JENCI	E DES	SCRIP'	TION:	SEQ	ID NO	:15:								
25		Ser	Asn	Arg	Phe	Ser 5	Gly	Ser	Lys	Ser	Gly 10	Asn	Thr	Ala	Ser	Leu 15
30		Thr	Ile	Ser	Gly	Leu 20	Gln	Ala	Glu	Asp	Glu 25	Ala	Asp	Tyr	Tyr	Cys 30
		Ser	Ser	Tyr	Thr	Thr 35	Arg	Ser	Thr	Arg	Val 40	Phe	Gly	Gly	Gly	Thr 45
35		Lys	Leu	Thr	Val	Leu 50		•				٠		· .		
40	(2) INFORMA	ATION	FOR S	SEQ II	O NO:	16:										
	(i) SEQU	ENCE	CHA	RACTI	ERIST	ICS:										
45	(B) T	ENGT YPE: <i>i</i> OPOL	Amino	Acid												
	(xi) SEQU	JENCI	E DES	CRIP	TION:	SEQ	ID NO	:16:								
50		Ser 1	Asn	Arg	Phe	Ser 5	Gly	Ser	Lys	Ser	Gly 10	Asn	Thr	Ala	Ser	Leu 15
55		Thr	Ile	Ser	Gly	Leu 20	Gln	Ala	Glu	Asp	Glu 25	Ala	Asp	Tyr	Tyr	Cys
55	·	Ser	Ser	Tyr	Thr	Thr	Arg	Ser	Thr	Arg	Val	Phe	Gly	Gly	Gly	Thr

						35					40				•	45
5		Lys	Leu	Thr	Val	Leu 50										
	(2) INFORMA	TION	FOR	SEQ	ID NO	:17:										
10	(i) SEQUE	ENCE	E CHA	RACT	ERIS	TICS:										
70	(B) T	YPE:	Amin	amin o Acid : Line		s										
15	(xi) SEQL	JENC	EDE	SCRIF	PTION	: SEQ	ID NO	D:17:								
20	\$	Ser 1	Asn	Arg	Phe	Ser 5	Gly	Ser	Lys	Ser	Gly 10	Asn	Thr	Ala	Ser	Leu 15
20	•	Thr	Ile	Ser	Gly	Leu 20	Gln	Ala	Glu	Asp	Glu 25	Ala	Asp	Tyr	Tyr	Cys 30
25	8	Ser	Ser	Tyr	Thr	Thr 35	Arg	Ser	Thr	Arg	Val 40	Phe	Gly	Gly	Gly	Thr 45
	1	Lys	Leu	Thr	Val	Leu 50										
30	(2) INFORMA	TION	FOR	SEQ	ID N0:	18:										
	(i) SEQUE	ENCE	E CHA	RACT	ERIS	TICS:										
<i>35</i>	(B) T	YPE:	Amin	o Acid		S										
	(D) T	OPOI	LOGY	': Line	ar											
	(xi) SEQL	JENC	EDE	SCRIF	PTION	: SEQ	ID NO	D:18:								
40	S	Ser 1	Asn	Arg _.	Phe	Ser 5	Gly	Ser	Lys	Ser	Gly 10	Asn	Thr	Ala	Ser	Leu 15
45	.	Chr	Ile	Ser	Gly	Leu 20	Gln	Ala	Glu	Asp	Glu 25	Ala	Asp	Tyr	Tyr	Cys 30
	S	Ser	Ser	Tyr	Thr	Thr 35	Arg	Ser	Thr	Arg	Val 40	Phe	Gly	Gly	Gly	Thr 45
50		Lys	Leu	Thr	Val	Leu 50										
	(2) INFORMA	TION	FOR	SEQ	ID NO	:19:										
<i>55</i>	(i) SEQUE	ENCE	E CHA	RACT	ERIS	TICS:										
	` '			amin o Acid	o acid	S										

(D) TOPOLOGY: Linear

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:															
5		Can	7.0	a Base	• 17h -	. Com	- ~7-	. 50-	. T		. (2) -		m \}	27.		
		ser 1		. Arg	3 bµe	s ser S		, ser	. Lys	s ser	10		unr	. ATS	. Ser	Lev 15
10		Thr	: Ile	e Sei	c Gly	Leu	ı Glı	n Ala	a Glı	ı Asp	Glu	Ala	Asp	Tyr	Tyr	Cys
						20					25					30
15		Ser	Ser	Tyr	Thr	Thr 35	Arg	Ser	Thr	Arg	Val 40	Phe	Gly	Gly	Gly	Thr 45
		Lys	Leu	Thr	Val	Leu 50				-						
20	(2) INFOR	RMATIO	N FOI	RSEC) ID NO	D:20:										
	(i) SE	QUENC	E CH	ARAC	TERIS	STICS:										
25	(I	A) LENG B) TYPE D) TOP(: Ami	no Aci	d	ds										
30	(xi) S	EQUEN	CE DI	ESCR	IPTION	N: SEC	A DI Ç	IO:20:								
		Ser 1	Asn	Arg	Phe	Ser 5	Gly	Ser	Lys	Xaa	Gly 10	Asn	Thr	Ala	Ser	Leu 15
35		Thr	Ile	Ser	Gly	Leu 20	Gln	Ala	Glu	qaA	Glu 25	Ala	Asp	Tyr	Tyr	Cys 30
40		Ser	Ser	Tyr	Thr	Thr 35	Arg	Ser	Thr	Arg	Val 40	Phe	Gly	Gly	Gly	Thr 45
40		Lys	Leu	Thr	Val	Leu 50	•			•			•	٠		
45	(2) INFOR	RMATIO	N FOI	RSEC) ID NO	D:21:										
	(i) SE	QUENC	E CH	ARAC	TERIS	STICS:	•									
50	(I	A) LENG B) TYPE D) TOP(: Ami	no Aci	d	ds										
	(xi) S	EQUEN	CE DI	ESCR	IPTION	N: SEC	A DI Ç	l0:21:								

		Ser 1	Asn	Arg	Phe	Ser 5	_	Ser	Lys	Ser	10 Gly	Asn	Thr	Ala	Ser	Leu 15
5		Thr	Ile	Ser	Gly	Leu 20		Ala	Glu	Asp	Glu 25	Ala	Asp	Tyr	Tyr	Cys 30
		Ser	Ser	Tyr	Thr	Thr 35		Ser	Thr	Arg	Val 40	Phe	Gly	Gly	Gly	Thr 45
10		Lys	Leu	Thr	Val	Leu 50										
15	(2) INFORM	MATION	N FOR	RSEQ	ID No	:22:										
	(i) SEQ	UENC	E CHA	ARAC	TERIS	TICS:	:									
20	(B)	LENG TYPE: TOPO	: Amir	no Acid	d	ls										
	(xi) SE	QUENC	CE DE	SCRI	PTION	I: SEC	N DI Ç	0:22:								
25		Ser	Asn	Arg	Phe	Ser	: Gly	ser Ser	: Lys	Ser	Gly	Asn	Thr	: Ala	Ser	Leu
		1				5					10					15
30			Ile	Ser	Gly	_	Gln	Ala	Glu	Asp		Ala	asp	Tyr	Tyr	
						20			•	~	25		•	• (•	30
35		Ser	Ser	Tyr	Thr	Thr 35	Arg	Ser	Thr	Arg	Val 40	Phe	Gly	Gly	Gly	Thr 45
		Lys	Leu	Thr	Val	Leu 50										
40	(2) INFORM	MATION	N FOR	R SEQ	ID No	:23:										
	(i) SEG	UENC	E CHA	ARAC	TERIS	TICS:	:									
45	(B)	LENG TYPE: TOPO	: Amir	no Acid	d	ids										
50	(xi) SE	QUENC	CE DE	ESCRI	PTION	I: SEC	N DI Ç	O:23:								
<i>55</i>																
<i>JJ</i>																

		Gln 1	Val	Gln	Leu	Val 5	Gln	Ser	Gly	Gly	Gly 10	Leu	Val	Gln	Pro	Gly 15
5		Gly	Ser	Leu	Arg	Leu 20	Ser	Cys	Ala	Ala	Ser 25	Gly	Phe	Thr	Phe	Ser 30
4.0		Ser	Tyr	Glu	Met	Asn 35	Trp	Val	Arg	Gln	Ala 40	Pro	Gly	Lys	Gly	Leu 45
10		Glu	Trp	Val	Ser	Gly 50	Ile	Ser	Gly	Ser	Gly 55	Gly	Ser	Thr	Tyr	Tyr 60
15		Ala	Asp	Ser	Val	Lys 65	Gly	Arg	Phe	Thr	Ile 70	Ser	Arg	qzA	Asn	Ser 75
		Lys	Asn	Thr	Leu	Tyr 80	Leu	Gln	Met	Asn	Arg 85	Leu	Arg	Ala	Glu	Asp 90
20		Thr	Ala	Val	Tyr	Tyr 95	Cys	Ala	Arg	Asp	Asn 100	Gly	Trp	Glu	Leu	Thr 105
		Asp	Trp	Tyr	Phe	Asp 110	Leu	Trp	Gly	Arg	Gly 115	Thr	Met	Val	Thr	Val 120
25		Ser	Ser 122				•									
30	(2) INFORM	ATION	I FOR	SEQ	ID NO	D:24:										
	(i) SEQL	JENCI	E CHA	ARAC	TERIS	STICS:										
<i>35</i>	(B) -	ΓΥΡΕ:	Amir	23 am io Acid /: Line		ids										
	(xi) SEQ	UENC	DE DE	SCRI	PTION	N: SEC	Q ID N	O:24:								
40		Glu 1	Val	Gln	Leu	Val		Ser	Gly	Pro	Gly		. Val	Lys	Pro	Ser 15
		Gln	Thr	Leu	Ser	Leu	Thr	Cys	Thr	· Val	Sex	Gly	Gly	Ser	· Ile	Ser
45																
50																
55																

						20					25					30
5	S	er	Gly	Gly	Tyr	Tyr 35	Trp	Ser	Trp	Ile	Arg 40	Gln	His	Pro	Gly	Lys 45
	G	Sly	Leu	Glu	Trp	Ile 50	Gly	Tyr	Ile	Tyr	Tyr 55	Ser	Gly	Ser	Thr	Tyr 60
10	T	yr	Asn	Pro	Ser	Leu 65	Lys	Ser	Arg	Val	Thr 70	Ile	Ser	Val	Asp	Thr 75
15	S	Ser	Lys	Asn	Gln	Phe 80	Ser	Leu	Lys	Leu	Ser 85	Ser	Val	Thr	Ala	Ala 90
	7	4sp	Thr	Ala	Val	Tyr 95	Tyr	Cys	Ala	Arg	Val 100	Asp	Leu	Glu	Asp	Tyr 105
20	G	Ely	Ser	Gly	Ala	Ser 110	Asp	Tyr	Trp	Gly	Gln 115	Gly	Thr	Leu	Val	Thr 120
	V	/al	Ser	Ser 123						•		•				
25	(2) INFORM	ATIC	ON FC	OR SE	Q ID N	10:25:										
	(i) SEQU	JEN	CE CI	HARA	CTER	ISTIC	S:									
30	(B)	TYP	E: Am	107 a nino Ad GY: Lir	cid	acids										
<i>35</i>	(xi) SEG)UEN	NCE [DESCF	RIPTIC	ON: SE	EQ ID	NO:25	5 :							
40																
45																
50																

	Asp 1	Ile	Gln	Met	Thr 5	Gln	Ser	Pro	Ser	Thr 10	Leu	Ser	Ala	Ser	Ile 15
5	Gly	Asp	Arg	Val	Thr 20	Ile	Thr	Cys	Arg	Ala 25	Ser	Glu	Gly	Ile	Tyr 30
10	His	Trp	Leu	Ala	Trp 35	Tyr	Gln	Gln	Lys	Pro 40	Gly	Lys	Ala	Pró	Lys 45
10	Leu	Leu	Ile	Tyr	Lys 50	Ala	Ser	Ser	Leu	Ala 55	Ser	Gly	Ala	Pro	Ser 60
15	Arg	Phe	Ser	Gly	Ser 65	Gly	Ser	Gly	Thr	Asp 70	Phe	Thr	Leu	Thr	Ile 75
	Ser	Ser	Leu	Gln	Pro 80	Asp	Asp	Phe	Ala	Thr 85	Tyr	Tyr	Cys	Gln	Gln 90
20	Түх	Ser	Asn	Tyr	Pro 95		Thr	Phe	Gly	Gly	Gly	Thr	Lys	Leu	Glu 105
	Ile	Lys 107		•											
25	(2) INFORMATIO	ON FO	R SEG	N DI Ç	O:26:										
	(i) SEQUEN	CE CH	HARAG	CTERI	STICS	3 :									
30	(A) LEN (B) TYP (D) TOP	E: Am	ino Ac	id	cids										
35	(xi) SEQUEN	NCE D	ESCF	RIPTIO	N: SE	Q ID N	NO:26:								
40															
45															
50															
<i>55</i>															

	Asn 1	Ala	Tyr	Ala	Leu 5	Lys	Met	Ala	Asp	Pro 10	Asn	Arg	Phe	Arg	Gly 15
5	Lys	Asp	Leu	Ala	Ala 20	His	Tyr	Gly	Gln	Pro 25	Arg	Glu	Pro	Gln	Val 30
10	Tyr	Thr	Leu	Pro	Pro 35	Ser	Arg	Glu	Glu	Met 40	Thr	Lys	Asn	Gln	Val 45
	Ser	Leu	Trp	Суѕ	Leu 50	Val	Lys	Gly	Phe	Tyr 55	Pro	Ser	Asp	Ile	Ala 60
15	Val	Glu	Trp	Glu	Ser 65	Asn	Gly	Gln	Pro	Glu 70	Asn	Asn	Tyr	Lys	Thr 75
20	Thr	Pro	Pro	Val	Leu 80	Asp	Ser	Asp	Gly	Ser 85	Phe	Phe ·	Leu	Tyr	Ser 90
			·		95	,	Ser	-		100		_			105
25		-			110					115					Gln . 120
					125		Pro		,	130					135
30					140		Met			145					150
35		_		_	155		Arg			160		-			165
		,			170		Pro			175				-	180
40					185		Asn		-	190				_	195
					200		Phe			205					210
45		-			215		Gln			220					225
50		_		_	230		Asn			235			_		240
			Pro		245				4 ~	250	-		~ ~ ~		255
55			- 	~ ~ I		261									

(2) INFORMATION FOR SEQ ID NO:27:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 717 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single

(D) TOPOLOGY: Linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

CAGGTGCAGC TGGTGCAGTC TGGGGGGAGGC TTGGTCCGGC CCGGGGGGTC 50 CCTGAGTCTC TCCTGTGCAG TCTCTGGAAT CACCCTCAGG ACCTACGGCA 100 15 TGCACTGGGT CCGCCAGGCT CCAGGCAAGG GGCTGGAGTG GGTGGCAGGT 150 ATATCCTTTG ACGGAAGAAG TGAATACTAT GCAGACTCCG TGCAGGGCCG 200 ATTCACCATC TCCAGAGACA GTTCCAAGAA CACCCTGTAT CTGCAAATGA 250 ACAGCCTGAG AGCCGAGGAC ACGGCTGTGT ATTACTGTGC GAGAGGAGCA 300 CATTATGGTT TCGATATCTG GGGCCAAGGG ACAATGGTCA CCGTCTCGAG 350 25 TGGTGGAGGC GGTACAGGCG GAGGTGGCAG CGGCGGTGGC GGATCGGACA 400 TCCAGATGAC CCAGTCTCCT TCCACCCTGT CTGCATCTAT TGGAGACAGA 450 GTCACCATCA CCTGCCGGGC CAGCGAGGGT ATTTATCACT GGTTGGCCTG 500 GTATCAGCAG AAGCCAGGGA AAGCCCCTAA ACTCCTGATC TATAAGGCCT 550 CTAGTTTAGC CAGTGGGGCC CCATCAAGGT TCAGCGGCAG TGGATCTGGG 600 ACAGATTICA CICTCACCAI CAGCAGCCIG CAGCCIGAIG ATTITIGCAAC 650 TTATTACTGC CAACAATATA GTAATTATCC GCTCACTTTC GGCGGAGGGA 700 CCAAGCTGGA GATCAAA 717

(2) INFORMATION FOR SEQ ID N0:28:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 732 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

55

5

10

20

30

35

40

45

	CAGGIGCAGC	IGGIGCAAIC	IGGGGGAGGC	TIGGIACAGC	HIGGHGGGIC	50
5	CCTGAGACTC	TCCTGTGCAG	CCTCTGGATT	CACCTTCAGT	AGTTATGAAA	100
	TGAACTGGGT	CCGCCAGGCT	CCAGGGAAGG	GGCTGGAGTG	GGTCTCAGGT	150
	ATTAGTGGTA	'GTGGTGGTAG	CACATACTAC	GCAGACTCCG	TGAAGGGCCG	200
10	GTTCACCATC	TCCAGAGACA	ATTCCAAGAA	CACGCTGTAT	CTGCAAATGA	250
	ACAGACTGAG	AGCTGAGGAC	ACGGCTGTGT	ATTACTGTGC	GAGAGATAAT	300
15	GGGTGGGAAC	TAACGGACTG	GTACTTCGAT	CTCTGGGGCC	GGGGGACAAT	350
	GGTCACCGTC	TCCTCAGGTG	GAGGCGGTTC	AGGCGGAGGT	GGCAGCGGCG	400
20	GTGGCGGATC	GGACATCCAG	ATGACCCAGT	CTCCTTCCAC	CCTGTCTGCA	450
	TCTATTGGAG	ACAGAGTCAC	CATCACCTGC	CGGGCCAGTG	AGGGTATTTA	500
	TCACTGGTTG	GCCTGGTATC	AGCAGAAGCC	AGGGAAAGCC	CCTAAACTCC	550
25	TGATCTATAA	GGCCTCTAGT	TTAGCCAGTG	GGGCCCCATC	AAGGTTCAGC	600
	GGCAGTGGAT	CTGGGACAGA	TTTCACTCTC	ACCATCAGCA	GCCTGCAGCC	650
30	TGATGATTTT	GCAACTTATT	ACTGCCAACA	ATATAGTAAT	TATCCGCTCA	700
	CTTTCGGCGG	AGGGACCAAG	CTGGAGATCA	AA 732		

35 Claims

40

45

50

55

1. A method of preparing a multispecific antibody comprising at least two different binding domains, wherein a first binding domain binds a first molecule and a second binding domain binds a second molecule different

from said first molecule, each binding domain comprises (i) a heavy chain variable domain which is part of a polypeptide which further

the first binding domain comprises a first heavy chain variable domain of a first polypeptide, the second binding domain comprises a second heavy chain variable domain of a second polypeptide, and the first and second heavy chain variable domains are different, and

the polypeptides multimerize by interaction of said multimerization domains, and wherein

comprises a multimerization domain and (ii) a variable domain of a light chain,

either each light chain variable domain is identical,

or the multispecific antibody comprises a first light chain variable domain and a second light chain variable domain, the first and second light chain variable domains differ from one another but have at least 80% amino acid sequence identity, and said first binding domain binds said first molecule whether said first binding domain comprises the first light chain variable domain or the second light chain variable domain, and said second binding domain binds said second molecule whether said second binding domain comprises the second light chain variable domain or the first light chain variable domain,

the method comprising the steps of:

(i) culturing a host cell comprising nucleic acid encoding said polypeptides and light chain or light chains, wherein the culturing is such that the nucleic acid is expressed and the polypeptides and light chain or light chains produced; and

(ii) recovering the multispecific antibody from the host cell culture.

5

10

25

- 2. The method of claim 1, wherein nucleic acid encoding the multimerization domain of said first polypeptide or nucleic acid encoding the multimerization domain of said second polypeptide, or both, is provided by alteration of nucleic acid to alter the encoded amino acid sequence.
- 3. The method of claim 2 wherein nucleic acid encoding the multimerization domain of said first polypeptide, said second polypeptide, or both, is altered so that the multimerization domain of each of said first or second polypeptide comprises a free thiol-containing residue which forms a disulfide bond with a free thiol-containing residue of the multimerization domain of the other of said first or second polypeptide.
- 4. The method of claim 1 wherein multimerization of said polypeptides comprises a protuberance-into-cavity interaction, wherein the method further comprises before step (i):
- providing nucleic acid encoding the first polypeptide by alteration of nucleic acid to generate a protuberance in the multimerization domain of the encoded polypeptide by replacing an amino acid residue with an import residue having a larger side chain volume, and providing nucleic acid encoding the second polypeptide by alteration of nucleic acid to generate a compensatory cavity in the multimerization domain of the encoded polypeptide by replacing an amino acid residue with an import residue having a smaller side chain volume.
 - 5. The method of claim 4 wherein nucleic acid is provided encoding a first polypeptide comprising a multimerization domain which has a protuberance, a second polypeptide comprising a multimerization domain which has a cavity, or a first polypeptide comprising a multimerization domain which has a protuberance and a second polypeptide comprising a multimerization domain which has a cavity, by means of phage display selection.
 - 6. The method of claim 4 wherein the import residue having a larger side chain volume is selected from the group consisting of arginine (R), phenylalanine (F), tyrosine (Y), (W), isoleucine (I) and leucine (L)
- 7. The method of claim 4 wherein the import residue having a smaller side chain volume is selected from the group consisting of glycine (G), alanine (A), serine (S), threonine (T), and valine (V), and wherein the import residue is not cysteine (C).
- 8. The method of any of claims 1-7, wherein the first and second polypeptide each comprise an antibody constant domain.
 - 9. The method of claim 8 wherein the first and second polypeptide each comprise an antibody constant domain selected from the group consisting of a C_H3 domain and an IgG constant domain.
- 10. The method of claim 1, wherein the multispecific antibody is an immunoadhesin.
 - 11. The method of claim 1, further comprising a step preceding step (i) wherein said nucleic acid is introduced into the host cell.
- 12. The method of any of claims 1-11, wherein each light chain variable domain is identical.
 - 13. The method of claim 1, wherein the multispecific antibody comprises a first light chain variable domain and a second light chain variable domain, and the first and second light chain variable domains differ from one another but have at least 90% amino acid sequence identity.
 - 14. The method of claim 13, wherein the first and second light chain variable domains have at least 95% amino acid sequence identity.
- **15.** The method of claim 14, wherein the first and second light chain variable domains have at least 98% sequence identity.
 - 16. The method of claim 15, wherein the first and second light chain variable domains have at least 99% sequence identity.

- 17. A multispecific antibody comprising at least two different binding domains, wherein
 - a first binding domain binds a first molecule and a second binding domain binds a second molecule different from said first molecule,
 - each binding domain comprises (i) a heavy chain variable domain which is part of a polypeptide which further comprises a multimerization domain and (ii) a variable domain of a light chain,

the first binding domain comprises a first heavy chain variable domain of a first polypeptide, the second binding domain comprises a second heavy chain variable domain of a second polypeptide, and the first and second heavy chain variable domains are different, and

the polypeptides multimerize by interaction of said multimerization domains, and wherein

either each light chain variable domain is identical,

5

10

15

20

25

or the multispecific antibody comprises a first light chain variable domain and a second light chain variable domain, the first and second light chain variable domains differ from one another but have at least 80% amino acid sequence identity, an said first binding domain binds said first molecule whether said first binding domain comprises the first light chain variable domain or the second light chain variable domain, and said second binding domain binds said second molecule whether said second binding domain comprises the second light chain variable domain or the first light chain variable domain.

- **18.** The multispecific antibody of claim 17, wherein the multimerization domain of the first polypeptide or multimerization domain of the second polypeptide, or both, is provided by alteration of an amino acid.
- 19. The multispecific antibody of claim 18 wherein the multimerization domain of each of said first or second polypeptide comprises free thiol-containing residue which forms a disulfide bond with a free thiol-containing residue of the multimerization domain of the other of said first or second polypeptide.
- 20. The multispecific antibody of claim 18 wherein multimerization of said polypeptides comprises a protuberance-into-cavity interaction and the multimerization domain of the first polypeptide comprises a protuberance and the multimerization domain of the second polypeptide comprises a compensatory cavity.
- 21. The multispecific antibody of claim 20 wherein the protuberance and cavity are generated by alterations in which naturally occurring amino acids are imported into the first and second polypeptides.
 - 22. The multispecific antibody of any of claims 17-21, wherein each light chain variable domain is identical.
- 23. The multispecific antibody of claim 17, wherein the multispecific antibody comprises a first light chain variable domain and a second light chain variable domain, and the first and second light chain variable domains differ from one another but have at least 90% amino acid sequence identity.
- 24. The multispecific antibody of claim 23, wherein the first and second light chain variable domains have at least 95% amino acid sequence identity.
 - 25. The multispecific antibody of claim 24, wherein the first and second light chain variable domains have at least 98% sequence identity.
- 26. The multispecific antibody of claim 25, wherein the first and second light chain variable domains have at least 99% sequence identity.
 - 27. A composition comprising the multispecific antibody of claim 17 and a carrier.
- 28. A host cell comprising nucleic acid encoding the multispecific antibody of claim 17.
 - 29. The host cell of claim 28 wherein the host cell is a mammalian cell.
- **30.** A method of preparing a multispecific antibody comprising a first binding domain which binds a first molecule and a second binding domain which binds a second molecule different from said first molecule, the method comprising:
 - (a) selecting a first nucleic acid encoding a first polypeptide comprising a first heavy chain variable domain and a multimerization domain, and selecting a second nucleic acid encoding a second polypeptide comprising

a second heavy chain variable domain and a multimerization domain, wherein the first and second heavy chain variable domains are different and the multimerization domain of each of said first and second polypeptides comprises an amino acid residue that specifically interacts with an amino acid residue in the multimerization domain of the other of said first and second polypeptides, thereby generating a stable interaction between the first and second polypeptides;

(b) either

5

10

15

20

30

35

- (i) selecting nucleic acid encoding a variable light chain, wherein the variable light chain interacts with each first and second polypeptide to form said first and second binding domains, or
- (ii) selecting a first variable light chain-encoding nucleic acid and a second variable light chain-encoding nucleic acid, wherein the first and second variable light chains differ from one another but have at least 80% amino acid sequence identity, each of the first and second variable light chains interacts with one of the first and second polypeptides to form said first and second binding domains, and said first binding domain binds said first molecule whether said first binding domain is formed by interaction of the first polypeptide with the second variable light chain, and said second binding domain binds said second molecule whether said second binding domain is formed by interaction of the second polypeptide with the second variable light chain or interaction of the second polypeptide with the first variable light chain;
- (c) introducing into a host cell the nucleic acid encoding the first and second polypeptides and the variable light chain or variable light chains, and culturing the cell so that expression of the nucleic acid occurs and the encoded polypeptides and variable light chain or variable light chains are produced;
- (d) recovering the multispecific antibody from the cell culture.
- 25 31. The method of claim 30, wherein said first nucleic acid encoding the first polypeptide, said second nucleic acid encoding the second polypeptide, or both, are selected from nucleic acid altered to alter the encoded amino acid sequence.
 - 32. The method of claim 31 wherein the first and second polypeptides interact by a protuberance-into-cavity interaction.
 - 33. The method of claim 31 wherein nucleic acid is altered to import a free thiol-containing residue into the encoded amino acid sequence.
 - 34. The method of claim 30, wherein the first and second polypeptide each comprise an antibody constant domain.
 - 35. The method of claim 34 wherein the antibody constant domain is a C_H 3 domain.
 - 36. The method of claim 34 wherein the antibody constant domain is from a human IgG.
- 37. The method of claim 30, wherein nucleic acid is selected in accordance with (i).
 - 38. The method of claim 30, wherein nucleic acid is selected in accordance with (ii).
 - **39.** The method of claim 38 wherein the first and second variable light chains differ from one another but have at least 90% amino acid sequence identity.
 - **40.** The method of claim 39, wherein the first and second variable light chains have at least 95% amino acid sequence identity.
- 50 **41.** The method of claim 40, wherein the first and second variable light chains have at least 98% sequence identity.
 - 42. The method of claim 41, wherein the first and second variable light chains have at least 99% sequence identity.
- 43. A method of measuring the formation of a heteromultimeric multispecific antibody from a mixture of polypeptides,

 wherein the multispecific antibody comprises at least two different binding domains, wherein
 - a first binding domain binds a first molecule and a second binding domain binds a second molecule different from said first molecule,
 - each binding domain comprises (i) a heavy chain variable domain which is part of a polypeptide which further

comprises a multimerization domain and (ii) a variable domain of a light chain,

the first binding domain comprises a first heavy chain variable domain of a first polypeptide, the second binding domain comprises a second heavy chain variable domain of a second polypeptide, and the first and second heavy chain variable domains are different, and

the polypeptides multimerize by interaction of said multimerization domains, wherein the multimerization domain of each of said first or second polypeptide comprises a free thiol-containing residue which forms a disulfide bond with a free thiol-containing residue of the multimerization domain of the other of said first or second polypeptide

and wherein

5

10

15

20

35

40

50

55

either each light chain variable domain is identical,

or the multispecific antibody comprises a first light chain variable domain and a second light chain variable domain, the first and second light chain variable domains differ from one another but have at least 80% sequence identity, and said first binding domain binds said first molecule whether said first binding domain comprises the first light chain variable domain or the second light chain variable domain, and said second binding domain binds said second molecule whether said second binding domain comprises the second light chain variable domain or the first light chain variable domain,

the method comprising the steps of:

- (a) causing each the multispecific antibodies to migrate in a gel matrix; and
- (b) determining the relative amount of a band corresponding to the multispecific antibody having a non-naturally occurring disulfide bond between the first and second polypeptides, and a slower migrating band corresponding to a heteromultimer lacking non-naturally occurring disulfide bonds between the first and second polypeptides.
- 25 44. The method of claim 43 wherein multimerization of said polypeptides is promoted by a protuberance-into-cavity interaction between the multimerization domains.
 - 45. The method of claim 43 wherein each light chain variable domain is identical.
- 46. The method of claim 43, wherein the first and second light chain variable domains differ from one another but have at least 90% amino acid sequence identity.
 - 47. The method of claim 46, wherein the first and second light chain variable domains have at least 95% amino acid sequence identity.
 - 48. The method of claim 47, wherein the first and second variable light claims have at least 98% sequence identity.
 - **49.** The method of claim 48, wherein the first and second light chain variable domains have at least 99% sequence identity.
 - **50.** The method according to claim 1, wherein the multispecific antibody is selected from the group consisting of antihuman leptin receptor ECD (Ob-R)/anti- human receptor tyrosine kinase HER3 and anti-human thrombopoietin receptor tyrosine kinase (Mpl)/anti-human receptor tyrosine kinase HER3.
- 51. The multispecific antibody of claim 17, which is selected from the group consisting of anti-human leptin receptor ECD (Ob-R)/anti-human receptor tyrosine kinase HER3 and anti-human thrombopoietin receptor tyrosine kinase (MpI)/anti-human receptor tyrosine kinase HER3.
 - 52. A composition comprising the multispecific antibody of claim 51 and a carrier.
 - **53.** The host cell of claim 28, wherein the multispecific antibody is selected from the group consisting of anti-human leptin receptor ECD (Ob-R)/anti-human receptor tyrosine kinase HER3 and anti-human thrombopoietin receptor tyrosine kinase (Mpl)/anti-human receptor tyrosine kinase HER3.

Patentansprüche

1. Verfahren zur Herstellung eines multispezifischen Antikörpers, umfassend zumindest zwei unterschiedliche Bin-

dedomänen, worin

eine erste Bindedomäne ein erstes Molekül bindet und eine zweite Bindedomäne ein zweites Molekül bindet, das sich vom ersten Molekül unterscheidet,

wobei jede Bindedomäne (i) eine variable Schwerketten-Domäne, die Teil eines Polypeptids ist, das weiters eine Multimerisations-Domäne umfasst, und (ii) eine variable Leichtketen-Domäne umfasst,

worin die erste Bindedomäne eine erste variable Schwerketten-Domäne eines ersten Polypeptids umfasst, die zweite Bindedomäne eine zweite variable Schwerketten-Domäne eines zweiten Polypeptids umfasst und sich die erste und die zweite variable Schwerketten-Domäne unterscheiden, und

die Polypeptide durch Wechselwirkung der Multimerisations-Domänen multimerisieren, und worin

entweder alle variablen Leichtketten-Domänen identisch sind,

oder der multispezifische Antikörper eine erste variable Leichtketten-Domäne und eine zweite variable Leichtketten-Domäne umfasst, worin sich die erste und die zweite variable Leichtketten-Domäne voneinander unterscheiden, jedoch zumindest 80 % Aminosäuresequenz-Identität aufweisen, und die erste Bindedomäne das erste Molekül bindet, unabhängig davon, ob die erste Bindedomäne die erste variable Leichtketten-Domäne umfasst, und die zweite Bindedomäne das zweite Molekül bindet, unabhängig davon, ob die zweite Bindedomäne die zweite variable Leichtketten-Domäne oder die erste variable Leichtketten-Domäne umfasst,

wobei das Verfahren folgende Schritte umfasst:

- (i) das Kultivieren einer Wirtszelle, die Nucleinsäure umfasst, die für die Polypeptide und die Leichtkette oder Leichtketten kodiert, worin das Kultivieren so erfolgt, dass die Nucleinsäure exprimiert wird und die Polypeptide und die Leichtkette oder Leichtketten produziert werden; und
- (ii) die Gewinnung des multispezifischen Antikörpers aus der Wirtszellkultur.
- 2. Verfahren nach Anspruch 1, worin für die Multimerisations-Domäne des ersten Polypeptids kodierende Nucleinsäure oder für die Multimerisations-Domäne des zweiten Polypeptids kodierende Nucleinsäure oder beide durch Änderung von Nucleinsäure zur Änderung der kodierten Aminosäuresequenz bereitgestellt wird/werden.
- 3. Verfahren nach Anspruch 2, worin Nucleinsäure, die für die Multimerisations-Domäne des ersten Polypeptids, des zweiten Polypeptids oder von beiden kodiert, so geändert wird, dass die Multimerisations-Domäne des ersten bzw. des zweiten Polypeptids jeweils einen freies Thiol enthaltenden Rest umfasst, der eine Disulfidbindung mit einem freies Thiol enthaltenden Rest der Multimerisations-Domäne des jeweils anderen, ersten oder zweiten, Polypeptids bildet.
 - 4. Verfahren nach Anspruch 1, worin Multimerisation des Polypeptids eine "Protuberanz-in-Hohlraum"- ("protuberance-in-cavity"-) Wechselwirkung umfasst, worin das Verfahren vor Schritt (i) weiters Folgendes umfasst:
 - das Bereitstellen von für das erste Polypeptid kodierender Nucleinsäure durch Änderung von Nucleinsäure zur Erzeugung einer Protuberanz in der Multimerisations-Domäne des kodierten Polypeptids durch Ersetzen eines Aminosäurerests durch einen Importrest mit einem größeren Seitenkettenvolumen, und das Bereitstellen von für das zweite Polypeptid kodierender Nucleinsäure durch Änderung von Nucleinsäure zur Erzeugung eines Kompensationshohlraums in der Multimerisations-Domäne des kodierten Polypeptids durch Ersetzen eines Aminosäurerests durch einen Importrest mit einem kleineren Seitenkettenvolumen.
 - 5. Verfahren nach Anspruch 4, worin Nucleinsäure, die für ein eine Multimerisations-Domäne mit einer Protuberanz umfassendes erstes Polypeptid, ein eine Multimerisations-Domäne mit einem Hohlraum umfassendes zweites Polypeptid, oder für ein eine Multimerisations-Domäne mit einer Protuberanz umfassendes erstes Polypeptid und ein eine Multimerisations-Domäne mit einem Hohlraum umfassendes zweites Polypeptid kodiert, mittels Phagendisplay-Selektion bereitgestellt wird.
 - 6. Verfahren nach Anspruch 4, worin der Importrest mit einem größeren Seitenkettenvolumen aus der aus Arginin (R), Phenylalanin (F), Tyrosin (Y), Tryptophan (W), Isoleucin (1) und Leucin (L) bestehenden Gruppe ausgewählt ist.
 - 7. Verfahren nach Anspruch 4, worin der Importrest mit einem kleineren Seitenkettenvolumen aus der aus Glycin (G), Alanin (A), Serin (S), Threonin (T) und Valin (V) bestehenden Gruppe ausgewählt ist.

20

25

5

10

15

30

35

45

40

55

- 8. Verfahren nach einem der Ansprüche 1 bis 7, worin das erste und das zweite Polypeptid jeweils eine konstante Antikörper-Domäne aufweisen.
- 9. Verfahren nach Anspruch 8, worin das erste und zweite Polypeptid jeweils eine konstante Antikörper-Domäne aufweisen, die aus der aus einer C_H3-Domäne und einer konstanten IgG-Domäne bestehenden Gruppe ausgewählt ist.
 - 10. Verfahren nach Anspruch 1, worin der multispezifische Antikörper ein Immunadhäsin ist.

5

30

35

40

45

50

- 11. Verfahren nach Anspruch 1, weiters umfassend einen Schritt vor Schritt (i), worin die Nucleinsäure in die Wirtszelle eingeführt wird.
 - 12. Verfahren nach einem der Ansprüche 1 bis 11, worin alle variablen Leichtketten-Domänen identisch sind.
- 13. Verfahren nach Anspruch 1, worin der multispezifische Antikörper eine erste variable Leichtketten-Domäne und eine zweite variable Leichtketten-Domäne umfasst und sich die erste und die zweite variable Leichtketten-Domäne voneinander unterscheiden, jedoch zumindest 90 % Aminosäuresequenz-Identität aufweisen.
- 14. Verfahren nach Anspruch 13, worin die erste und die zweite variable Leichtketten-Domäne zumindest 95 % Aminosäuresequenzidentität aufweisen.
 - 15. Verfahren nach Anspruch 14, worin die erste und die zweite variable Leichtketten-Domäne zumindest 98 % Sequenzidentität aufweisen.
- 16. Verfahren nach Anspruch 15, worin die erste und die zweite variable Leichtketten-Domäne zumindest 99 % Sequenzidentität aufweisen.
 - 17. Multispezifischer Antikörper, umfassend zumindest zwei unterschiedliche Bindedomänen, worin eine erste Bindedomäne ein erstes Molekül bindet und eine zweite Bindedomäne ein zweites Molekül bindet,

das sich vom ersten Molekül unterscheidet,

worin jede Bindedomäne (i) eine variable Schwerketten-Domäne, die Teil eines Polypeptids ist, das weiters eine Multimerisationsdomäne umfasst, und (ii) eine variable Leichtketten-Domäne umfasst,

worin die erste Bindedomäne eine erste variable Schwerketten-Domäne eines ersten Polypeptids, die zweite Bindedomäne eine zweite variable Schwerketten-Domäne eines zweiten Polypeptids umfasst und sich die erste und die zweite variable Schwerketten-Domäne unterscheiden, und

die Polypeptide durch Wechselwirkung der Multimerisations-Domänen multimerisieren, und worin

entweder alle variablen Leichtketten-Domänen identisch sind,

oder der multispezifische Antikörper eine erste variable Leichtketten-Domäne und eine zweite variable Leichtketten-Domäne umfasst, worin sich die erste und die zweite variable Leichtketten-Domäne voneinander unterscheiden, jedoch zumindest 80 % Aminosäuresequenz-Identität aufweisen, und die erste Bindedomäne das erste Molekül bindet, unabhängig davon, ob die erste Bindedomäne die erste variable Leichtketten-Domäne oder die zweite variable Leichtketten-Domäne umfasst, und die zweite Bindedomäne das zweite Molekül bindet, unabhängig davon, ob die zweite Bindedomäne die zweite variable Leichtketten-Domäne oder die erste variable Leichtketten-Domäne umfasst.

- 18. Multispezifischer Antikörper nach Anspruch 17, worin die Multimerisations-Domäne des ersten Polypeptids oder die Multimerisationsdomäne des zweiten Polypeptids oder beide durch Änderung einer Aminosäure bereitgestellt sind.
- 19. Multispezifischer Antikörper nach Anspruch 18, worin die Multimerisations-Domäne des ersten bzw. des zweiten Polypeptids jeweils einen freies Thiol enthaltenden Rest umfasst, der mit einem freies Thiol enthaltenden Rest der Multimerisationsdomäne des jeweils anderen, ersten oder zweiten, Polypeptids eine Disulfidbindung bildet.
- 20. Multispezifischer Antikörper nach Anspruch 18, worin Multimerisation der Polypeptide eine "Protuberanz-in-Hohl-raum"- ("protuberance-in-cavity"-) Wechselwirkung umfasst und die Multimerisations-Domäne des ersten Polypeptids eine Protuberanz und die Multimerisations-Domäne des zweiten Polypeptids einen Kompensationshohl-raum umfasst.

- 21. Multispezifischer Antikörper nach Anspruch 20, worin die Protuberanz und der Hohlraum durch Änderungen erzeugt sind, bei denen natürlich vorkommende Aminosäuren in das erste und das zweite Polypeptid importiert sind.
- 22. Multispezifischer Antikörper nach einem der Ansprüche 17 bis 21, worin alle variablen Leichtketten-Domänen identisch sind.
 - 23. Multispezifischer Antikörper nach Anspruch 17, worin der multispezifische Antikörper eine erste variable Leichtketten-Domäne umfasst, und sich die erste und die zweite variable Leichtketten-Domäne voneinander unterscheiden, jedoch zumindest 90 % Aminosäuresequenz-Identität aufweisen.
 - 24. Multispezifischer Antikörper nach Anspruch 23, worin die erste und die zweite variable Leichtketten-Domäne zumindest 95 % Aminosäuresequenz-Identität aufweisen.
- 25. Multispezifischer Antikörper nach Anspruch 24, worin die erste und die zweite variable Leichtketten-Domäne zumindest 98 % Sequenzidentität aufweisen.
 - 26. Multispezifischer Antikörper nach Anspruch 25, worin die erste und die zweite variable Leichtketten-Domäne zumindest 99 % Sequenzidentität aufweisen.
 - 27. Zusammensetzung, die einen multispezifischen Antikörper nach Anspruch 17 und einen Träger umfasst.
 - 28. Wirtszelle, die Nucleinsäure umfasst, die für einen multispezifischen Antikörper nach Anspruch 17 kodiert.
- 25 29. Wirtszelle nach Anspruch 28, worin die Wirtszelle eine Säugetierzelle ist.
 - **30.** Verfahren zur Herstellung eines multispezifischen Antikörpers, der eine erste Bindedomäne, die ein erstes Molekül bindet, und eine zweite Bindedomäne, die ein zweites Molekül bindet, das sich vom ersten Molekül unterscheidet, umfasst, worin das Verfahren Folgendes umfasst:
 - (a) das Auswählen einer ersten Nucleinsäure, die für ein eine erste variable Schwerketten-Domäne und eine Multimerisations-Domäne umfassendes erstes Polypeptid kodiert, und Auswählen einer zweiten Nucleinsäure, die für ein eine zweite variable Schwerketten-Domäne und eine Multimerisations-Domäne umfassendes zweites Polypeptid kodiert, worin sich die erste und die zweite variable Schwerketten-Domäne unterscheiden und die Multimerisations-Domäne des ersten und des zweiten Polypeptids jeweils einen Aminosäurerest umfassen, der spezifisch mit einem Aminosäurerest in der Multimerisations-Domäne des jeweils anderen, ersten oder zweiten, Polypeptids wechselwirkt, wodurch eine stabile Wechselwirkung zwischen dem ersten und dem zweiten Polypeptid erzeugt wird;
 - (b) entweder

5

10

20

30

35

40

45

50

- (i) das Auswählen von Nucleinsäure, die für eine variable Leichtkette kodiert, worin die variable Leichtkette sowohl mit dem ersten als auch mit dem zweiten Polypeptid wechselwirkt, um die erste und zweite Bindedomäne zu bilden.
 - oder
- (ii) das Auswählen von für die erste variable Leichtkette kodierender Nucleinsäure und von für die zweite variable Leichtkette kodierender Nucleinsäure, worin sich die erste und die zweite variable Leichtkette voneinander unterscheiden, jedoch zumindest 80 % Aminosäuresequenz-Identität aufweisen, die erste und die zweiten variable Leichtkette jeweils mit einem von erstem und zweitem Polypeptid wechselwirken, um die erste und die zweite Bindedomäne zu bilden, und die erste Bindedomäne das erste Molekül bindet, unabhängig davon, ob die erste Bindedomäne durch Wechselwirkung des ersten Polypeptids mit der zweiten variablen Leichtkette gebildet ist, und die zweite Bindedomäne das zweite Molekül bindet, unabhängig davon, ob die zweite Bindedomäne durch Wechselwirkung des zweiten Polypeptids mit der zweiten variablen Leichtkette oder durch Wechselwirkung des zweiten Polypeptids mit der zweiten variablen Leichtkette oder durch Wechselwirkung des zweiten Polypeptids mit der zweiten variablen Leichtkette gebildet ist;
- (c) das Einführen der für das erste und das zweite Polypeptid und die variable Leichtkette oder die variablen Leichtketten kodierendern Nucleinsäure in eine Wirtszelle und das Kultivieren der Zelle, sodass Expression der Nucleinsäure erfolgt und die kodierten Polypeptide und die variable Leichtkette oder die variablen Leicht-

ketten produziert werden;

5

10

20

30

35

40

45

50

55

- (d) die Gewinnung des multispezifischen Antikörpers aus der Zellkultur.
- 31. Verfahren nach Anspruch 30, worin die erste, für das erste Polypeptid kodierende Nucleinsäure, die zweite, für das zweite Polypeptid kodierende Nucleinsäure oder beide aus Nucleinsäuren ausgewählt werden, die zur Änderung der kodierten Aminosäuresequenz geändert werden.
- 32. Verfahren nach Anspruch 31, worin das erste und das zweite Polypeptid durch eine "Protuberanz-in-Hohlraum"- ("protuberance-in-cavity"-) Wechselwirkung wechselwirken.
- 33. Verfahren nach Anspruch 31, worin Nucleinsäure zum Import eines freies Thiol enthaltenden Rests in die kodierte Aminosäuresequenz geändert wird.
- **34.** Verfahren nach Anspruch 30, worin das erste und das zweite Polypeptid jeweils eine konstante Antikörper-Domäne umfassen.
 - 35. Verfahren nach Anspruch 34, worin die konstante Antikörper-Domäne eine C_H3-Domäne ist.
 - 36. Verfahren nach Anspruch 34, worin die konstante Antikörper-Domäne von menschlichem IgG herrührt.
 - 37. Verfahren nach Anspruch 30, worin Nucleinsäure gemäß (i) ausgewählt wird.
 - 38. Verfahren nach Anspruch 30, worin Nucleinsäure gemäß (ii) ausgewählt wird.
- 25 39. Verfahren nach Anspruch 38, worin sich die erste und die zweite variable Leichtkette voneinander unterscheiden, jedoch zumindest 90 % Aminosäuresequenz-Identität aufweisen.
 - **40.** Verfahren nach Anspruch 39, worin die erste und die zweite variable Leichtkette zumindest 95 % Aminosäuresequenz-Identität aufweisen.
 - **41.** Verfahren nach Anspruch 40, worin die erste und die zweite variable Leichtkette zumindest 98 % Sequenzidentität aufweisen.
 - **42.** Verfahren nach Anspruch 41, worin die erste und die zweite variable Leichtkette zumindest 99 % Sequenzidentität aufweisen.
 - 43. Verfahren zum Messen der Bildung eines heteromultimeren multispezifischen Antikörpers aus einem Polypeptid-Gemisch, worin der multispezifische Antikörper zumindest zwei unterschiedliche Bindedomänen umfasst, worin eine erste Bindedomäne ein erstes Molekül bindet und eine zweite Bindedomäne ein zweites Molekül bindet, das sich vom ersten Molekül unterscheidet,

jede Bindedomäne (i) eine variable Schwerketten-Domäne, die Teil eines Polypeptids ist, das weiters eine Multimerisations-Domäne umfasst, und (ii) eine variable Leichtketten-Domäne umfasst,

worin die erste Bindedomäne eine erste variable Schwerketten-Domäne eines ersten Polypeptids, die zweite Bindedomäne eine zweite variable Schwerketten-Domäne eines zweiten Polypeptids umfasst und sich die erste und die zweite variable Schwerketten-Domäne unterscheiden, und

die Polypeptide durch Wechselwirkung der Multimerisations-Domänen multimerisieren, worin die Multimerisationsdomäne des ersten oder des zweiten Polypeptids einen freies Thiol enthaltenden Rest umfasst, der mit einem freies Thiol enthaltenden Rest der Multimerisations-Domäne des jeweils anderen, ersten oder zweiten, Polypeptids eine Disulfidbindung bildet,

und worin

entweder alle variablen Leichtketten-Domänen identisch sind,

oder der multispezifische Antikörper eine erste variable Leichtketten-Domäne und eine zweite variable Leichtketten-Domäne umfasst, worin sich die erste und die zweite variable Leichtketten-Domäne voneinander unterscheiden, jedoch zumindest 80 % Aminosäuresequenz-Identität aufweisen, und die erste Bindedomäne das erste Molekül bindet, unabhängig davon, ob die erste Bindedomäne die erste variable Leichtketten-Domäne umfasst, und die zweite Bindedomäne das zweite Molekül bindet, unabhängig davon, ob die zweite Bindedomäne die zweite variable Leichtketten-Domäne oder die erste variable Leichtketten-Domäne umfasst,

wobei das Verfahren die folgenden Schritte umfasst:

- (a) das Auslösen von Migration beider multispezifischer Antikörper in eine Gelmatrix; und
- (b) das Bestimmen der relativen Menge einer Bande, die dem multispezifischen Antikörper mit einer nicht natürlich vorkommenden Disulfidbindung zwischen erstem und zweitem Polypeptid entspricht, und einer langsamer migrierenden Bande, die einem Heteromultimer ohne nicht natürlich vorkommende Disulfidbindung zwischen erstem und zweitem Polypeptid entspricht.
- 44. Verfahren nach Anspruch 43, worin Multimerisation der Polypeptide durch eine "Protuberanz-in-Hohlraum"- ("protuberance-in-cavity"-) Wechselwirkung zwischen den Multimerisations-Domänen unterstützt wird.
- 45. Verfahren nach Anspruch 43, worin alle variablen Leichtketten-Domänen identisch sind.
- **46.** Verfahren nach Anspruch 43, worin sich die erste und die zweite variable Leichtketten-Domäne voneinander unterscheiden, jedoch zumindest 90 % Aminosäuresequenz-tdentität aufweisen.
 - 47. Verfahren nach Anspruch 46, worin die erste und die zweite variable Leichtketten-Domäne zumindest 95 % Aminosäuresequenz-Identität aufweisen.
- 48. Verfahren nach Anspruch 47, worin die erste und die zweite variable Leichtketten-Domäne zumindest 98 % Sequenzidentität aufweisen.
 - **49.** Verfahren nach Anspruch 48, worin die erste und die zweite variable Leichtketten-Domäne zumindest 99 % Aminosäuresequenz-Identität aufweisen.
 - **50.** Verfahren nach Anspruch 1, worin der multispezifische Antikörper aus der aus Anti-Human-Leptin-Rezeptor ECD (Ob-R) / Anti-Human-Rezeptor-Tyrosin-Kinase HER3 und Anti-Human-Thrombopoietin-Rezeptor-Tyrosin-Kinase (MpI) / Anti-Human-Rezeptor-Tyrosin-Kinase HER3 bestehenden Gruppe ausgewählt ist.
- 51. Multispezifischer Antikörper nach Anspruch 17, der aus der aus Anti-Human-Leptin-Rezeptor ECD (Ob-R) / Anti-Human-Rezeptor-Tyrosin-Kinase HER3 und Anti-Human-Thrombopoietin-Rezeptor-Tyrosin-Kinase (Mpl) / Anti-Human-Rezeptor-Tyrosin-Kinase HER3 bestehenden Gruppe ausgewählt ist.
 - 52. Zusammensetzung, die einen multispezifischen Antikörper nach Anspruch 51 und einen Träger umfasst.
 - 53. Wirtszelle nach Anspruch 28, worin der multispezifische Antikörper aus der aus Anti-Human-Leptin-Rezeptor ECD (Ob-R) / Anti-Human-Rezeptor-Tyrosin-Kinase HER3 und Anti-Human-Thrombopoietin-Rezeptor-Tyrosin-Kinase (MpI) / Anti-Human-Rezeptor-Tyrosin-Kinase HER3 bestehenden Gruppe ausgewählt ist.

Revendications

5

10

15

25

35

40

45

50

- Méthode de préparation d'un anticorps multispécifique comprenant au moins deux domaines différents de liaison, où
 - un premier domaine de liaison lie une première molécule et un second domaine de liaison lie une seconde molécule différente de ladite première molécule,
 - chaque domaine de liaison comprend (i) un domaine variable d'une chaîne lourde qui fait partie d'un polypeptide qui comprend de plus un domaine de multimérisation et (ii) un domaine variable d'une chaîne légère,
 - le premier domaine de liaison comprend un premier domaine variable de chaîne lourde d'un premier polypeptide, le second domaine de liaison comprend un second domaine variable de chaîne lourde d'un second polypeptide et les premier et second domaines variables de chaîne lourde sont différents, et
 - les polypeptides se multimérisent par interaction desdits domaines de multimérisation,
 - et où soit chaque domaine variable de chaîne légère est identique
 - ou bien l'anticorps multispécifique comprend un premier domaine variable de chaîne légère et un second domaine variable de chaîne légère, les premier et second domaines variables de chaîne légère diffèrent l'un de l'autre mais ont au moins 80% d'identité des séquences d'acides aminés et ledit premier domaine de liaison lie ladite première molécule, que ledit premier domaine de liaison comprenne le premier domaine variable de chaîne légère ou le second domaine variable de chaîne légère et ledit second domaine de liaison lie ladite seconde

molécule, que ledit second domaine de liaison comprenne le second domaine variable de chaîne légère ou le premier domaine variable de chaîne légère,

la méthode comprenant les étapes de:

5

15

25

- (i) cultiver une cellule hôte comprenant un acide nucléique codant pour lesdits polypeptides et la chaîne légère ou les chaînes légères, où la culture est telle que l'acide nucléique soit exprimé et les polypeptides et la chaîne légère ou les chaînes légères produits; et
- (ii) récupérer l'anticorps multispécifique de la culture des cellules hôtes.
- 2. Méthode de la revendication 1, où l'acide nucléique codant pour le domaine de multimérisation dudit premier polypeptide ou l'acide nucléique codant pour le domaine de multimérisation dudit second polypeptide, ou les deux, est produit par altération de l'acide nucléique pour altérer la séquence codée d'acides aminés.
 - 3. Méthode de la revendication 2, où l'acide nucléique codant pour le domaine de multimérisation dudit premier polypeptide, dudit second polypeptide, ou des deux, est altéré de façon que le domaine de multimérisation de chacun desdits premier ou second polypeptides comprenne un résidu contenant un thiol libre qui forme une liaison disulfure avec un résidu contenant un thiol libre du domaine de multimérisation de l'autre dudit premier ou second polypeptide.
- 4. Méthode de la revendication 1, où la multimérisation desdits polypeptides comprend une interaction protubérancedans-cavité, où la méthode comprend de plus avant l'étape (i):
 - produire un acide nucléique codant pour le premier polypeptide par altération de l'acide nucléique pour générer une protubérance dans le domaine de multimérisation du polypeptide codé par remplacement d'un résidu d'acide aminé par un résidu d'import ayant un plus grand volume de chaîne latérale, et produire un acide nucléique codant pour le second polypeptide par altération de l'acide nucléique pour générer une cavité compensatoire dans le domaine de multimérisation du polypeptide codé par remplacement d'un résidu d'acide aminé par un résidu d'import ayant un plus petit volume de chaîne latérale.
- 5. Méthode de la revendication 4, où l'acide nucléique est produit codant pour un premier polypeptide comprenant un domaine de multimérisation qui a une protubérance, un second polypeptide comprenant un domaine de multimérisation qui a une cavité, ou un premier polypeptide comprenant un domaine de multimérisation qui a une protubérance et un second polypeptide comprenant un domaine de multimérisation qui a une cavité, au moyen d'une sélection de présentation de phage.
 - 6. Méthode de la revendication 4, où le résidu d'import ayant un plus grand volume de chaîne latérale est sélectionné dans le groupe consistant en arginine (R), phénylalanine (F), tyrosine (Y), tryptophane (W), isoleucine (I) et leucine (L).
- 7. Méthode de la revendication 4 où le résidu d'import ayant un plus petit volume de chaîne latérale est sélectionné dans le groupe consistant en glycine (G), alanine (A), sérine (S), thréonine (T), et valine (V), et où le résidu d'import n'est pas la cystéine (C).
- 8. Méthode de l'une quelconque des revendications 1-7 où les premier et second polypeptides forment chacun un domaine constant d'anticorps.
 - 9. Méthode de la revendication 8 où les premier et second polypeptides forment chacun un domaine constant d'anticorps sélectionné dans le groupe consistant en un domaine de C_H3 et un domaine constant de lgG.
- 10. Méthode de la revendication 1, où l'anticorps multispécifique est une immunoadhésine.
 - 11. Méthode de la revendication 1, comprenant de plus une étape précédant l'étape (i) où ledit acide nucléique est introduit dans la cellule hôte.
- 12. Méthode de l'une quelconque des revendications 1-11, où chaque domaine variable de chaîne légère est identique.
 - 13. Méthode de la revendication 1, où l'anticorps multispécifique comprend un premier domaine variable de chaîne légère et un second domaine variable de chaîne légère et les premier et second domaines variables de chaîne

légère diffèrent l'un de l'autre mais ont au moins 90% d'identité des séquences d'acides aminés.

- 14. Méthode de la revendication 13, où les premier et second domaines variables de chaîne légère ont au moins 95% d'identité des séquences d'acides aminés.
- 15. Méthode de la revendication 14, où les premier et second domaines variables de chaîne légère ont au moins 98% d'identité des séquences.
- 16. Méthode de la revendication 15, où les premier et second domaines variables de chaîne légère ont au moins 99% d'identité des séquences.
- 17. Anticorps multispécifique comprenant au moins deux domaines différents de liaison, où

un premier domaine de liaison lie une première molécule et un second domaine de liaison lie une seconde molécule différente de ladite première molécule,

chaque domaine de liaison comprend (i) un domaine variable de chaîne lourde qui fait partie d'un polypeptide qui comprend de plus un domaine de multimérisation et (ii) un domaine variable d'une chaîne légère,

le premier domaine de liaison comprend un premier domaine variable de chaîne lourde d'un premier polypeptide, le second domaine de liaison comprend un second domaine variable de chaîne lourde d'un second polypeptide et les premier et second domaines variables de chaîne lourde sont différents, et

les polypeptides se multimérisent par interaction desdits domaines de multimérisation,

et où

soit chaque domaine variable de chaîne légère est identique,

ou bien l'anticorps multispécifique comprend un premier domaine variable de chaîne légère et un second domaine variable de chaîne légère, les premier et second domaines variables de chaîne légère diffèrent l'un de l'autre mais ont au moins 80% d'identité des séquences d'acides aminés, un dit premier domaine de liaison lie ladite première molécule, que ledit premier domaine de liaison comprenne le premier domaine variable de chaîne légère ou le second domaine variable de chaîne légère et ledit second domaine de liaison lie ladite seconde molécule, que ledit second domaine de liaison comprenne le second domaine variable de chaîne légère ou le premier domaine variable de chaîne légère.

30

40

50

5

10

15

20

- 18. Anticorps multispécifique de la revendication 17, où le domaine de multimérisation du premier polypeptide ou le domaine de multimérisation du second polypeptide, ou les deux, est produit par altération d'un acide aminé.
- 19. Anticorps multispécifique de la revendication 18, où le domaine de multimérisation de chacun dudit premier ou second polypeptide comprend un résidu contenant un thiol libre qui forme une liaison disulfure avec un résidu contenant un thiol libre du domaine de multimérisation de l'autre dudit premier ou second polypeptide.
 - 20. Anticorps multispécifique de la revendication 18 où la multimérisation desdits polypeptides comprend une interaction protubérance-dans-cavité et le domaine de multimérisation du premier polypeptide comprend une protubérance et le domaine de multimérisation du second polypeptide comprend une cavité compensatoire.
 - 21. Anticorps multispécifique de la revendication 20, où la protubérance et la cavité sont générées par des altérations dans lesquelles des acides aminés naturels sont importés dans les premier et second polypeptides.
- 22. Anticorps multispécifique de l'une quelconque des revendications 17-21, où chaque domaine variable de chaîne légère est identique.
 - 23. Anticorps multispécifique de la revendication 17, où l'anticorps multispécifique comprend un premier domaine variable de chaîne légère et un second domaine variable de chaîne légère et les premier et second domaines variables de chaîne légère diffèrent l'un de l'autre mais ont au moins 90% d'identité des séquences d'acides aminés.
 - 24. Anticorps multispécifique de la revendication 23, où les premier et second domaines variables de chaîne légère ont au moins 95% d'identité des séquences d'acides aminés.
- 25. Anticorps multispécifique de la revendication 24, où les premier et second domaines variables de chaîne légère ont au moins 98% d'identité des séquences.
 - 26. Anticorps multispécifique de la revendication 24, où les premier et second domaines variables de chaîne légère

ont au moins 99% d'identité des séquences.

- 27. Composition comprenant l'anticorps multispécifique de la revendication 17 et un support.
- 5 28. Cellule hôte comprenant l'acide nucléique codant pour l'anticorps multispécifique de la revendication 17.
 - 29. Cellule hôte de la revendication 28, où la cellule hôte est une cellule mammalienne.
- 30. Méthode de préparation d'un anticorps multispécifique comprenant un premier domaine de liaison qui lie une première molécule et un second domaine de liaison qui lie une seconde molécule différente de ladite première molécule, la méthode comprenant:
 - (a) la sélection d'un premier acide nucléique codant pour un premier polypeptide comprenant un premier domaine variable de chaîne lourde et un domaine de multimérisation et la sélection d'un second acide nucléique codant pour un second polypeptide comprenant un second domaine variable de chaîne lourde et un domaine de multimérisation, où les premier et second domaines variables de chaîne lourde sont différents et le domaine de multimérisation de chacun desdits premier et second polypeptides comprend un résidu d'acide aminé qui interagit spécifiquement avec un résidu d'acide aminé dans le domaine de multimérisation de l'autre desdits premier et second polypeptides, pour ainsi générer une interaction stable entre les premier et second polypeptides;
 - (b) soit

15

20

25

30

35

40

45

- (i) la sélection d'un acide nucléique codant pour une chaîne légère variable, où la chaîne légère variable interagit avec chaque premier et second polypeptide pour former lesdits premier et second domaines de liaison, ou
- (ii) la sélection d'un acide nucléique codant pour une première chaîne légère variable et un acide nucléique codant pour une seconde chaîne légère variable, où les première et seconde chaînes légères variables diffèrent l'une de l'autre mais ont au moins 80% d'identité des séquences d'acides aminés, chacune des première et seconde chaînes légères variables interagit avec l'un des premier et second polypeptides pour former lesdits premier et second domaines de liaison et ledit premier domaine de liaison lie ladite première molécule, que ledit premier domaine de liaison soit formé par interaction du premier polypeptide avec la seconde chaîne légère variable, et ledit second domaine de liaison lie ladite seconde molécule, que ledit second domaine de liaison soit formé par interaction du second polypeptide avec la seconde chaîne légère variable ou interaction du second polypeptide avec la seconde chaîne légère variable ou interaction du second polypeptide avec la seconde chaîne légère variable;
- (c) l'introduction dans une cellule hôte de l'acide nucléique codant pour les premier et second polypeptides et la chaîne légère variable ou les chaînes légères variables, et la culture de la cellule de façon que l'expression de l'acide nucléique se produise et que les polypeptides codés et la chaîne légère variable ou les chaînes légères variables soient produits;
- (d) la récupération de l'anticorps multispécifique de la culture des cellules.
- **31.** Méthode de la revendication 30, où ledit premier acide nucléique codant pour le premier polypeptide, ledit second acide nucléique codant pour le second polypeptide, ou les deux, sont sélectionnés parmi un acide nucléique altéré pour altérer la séquence codée d'acides aminés.
- **32.** Méthode de la revendication 31, où les premier et second polypeptides interagissent par une interaction protubérance dans cavité.
- 33. Méthode de la revendication 31, où l'acide nucléique est altéré pour importer un résidu contenant un thiol libre dans la séquence d'acides aminés codés.
 - **34.** Méthode de la revendication 30, où les premier et second polypeptides comprennent chacun un domaine constant d'anticorps.
 - 35. Méthode de la revendication 34, où le domaine constant d'anticorps est un domaine de C_H3.
 - 36. Méthode de la revendication 34, où le domaine constant d'anticorps est de IgG humaine.

- 37. Méthode de la revendication 30, où l'acide nucléique est sélectionné selon (i).
- 38. Méthode de la revendication 30, où l'acide nucléique est sélectionné selon (ii).

10

15

20

25

30

35

40

45

- 39. Méthode de la revendication 38, où les première et seconde chaînes légères variables diffèrent l'une de l'autre mais ont au moins 90% d'identité des séquences d'acides aminés.
 - **40.** Méthode de la revendication 39, où les première et seconde chaînes légères variables ont au moins 95% d'identité des séquences d'acides aminés.
 - **41.** Méthode de la revendication 40, où les première et seconde chaînes légères variables ont au moins 98% d'identité des séquences.
 - **42.** Méthode de la revendication 41, où les première et seconde chaînes légères variables ont au moins 99% d'identité des séquences.
 - 43. Méthode de mesure de la formation d'un anticorps multispécifique hétéromultimère à partir d'un mélange de polypeptides, où l'anticorps multispécifique comprend au moins deux domaines différents de liaison, où

un premier domaine de liaison lie une première molécule et un second domaine de liaison lie une seconde molécule différente de ladite première molécule,

chaque domaine de liaison comprend (i) un domaine variable de chaîne lourde qui fait partie d'un polypeptide qui comprend de plus un domaine de multimérisation et (ii) un domaine variable d'une chaîne légère,

le premier domaine de liaison comprend un premier domaine variable de chaîne lourde d'un premier polypeptide, le second domaine de liaison comprend un second domaine variable de chaîne lourde d'un second polypeptide et les premier et second domaines variables de chaîne lourde sont différents; et

les polypeptides se multimérisent par interaction desdits domaines de multimérisation, où le domaine de multimérisation de chacun desdits premier ou second polypeptides comprend un résidu contenant un thiol libre qui forme une liaison disulfure avec un résidu contenant un thiol libre du domaine de multimérisation de l'autre desdits premier ou second polypeptides et où

soit chaque domaine variable de chaîne légère est identique,

ou l'anticorps multispécifique comprend un premier domaine variable de chaîne légère et un second domaine variable de chaîne légère, les premier et second domaines variables de chaîne légère diffèrent l'un de l'autre mais ont au moins 90% d'identité des séquences et ledit premier domaine de liaison lie ladite première molécule, que ledit premier domaine de liaison comprenne ledit premier domaine variable de chaîne légère ou ledit second domaine variable de chaîne légère et ledit second domaine de liaison lie ladite seconde molécule, que ledit second domaine de liaison comprennent le second domaine variable de chaîne légère ou le premier domaine variable de chaîne légère,

la méthode comprenant les étapes de:

- (a) forcer chacun des anticorps multispécifiques à migrer dans une matrice de gel; et
- (b) déterminer la quantité relative d'une bande correspondant à l'anticorps multispécifique ayant une liaison disulfure non naturelle entre les premier et second polypeptides et une bande migrant plus lentement correspondant à un hétéromultimère manquant de liaisons disulfures non naturelles entre les premier et second polypeptides.
- **44.** Méthode de la revendication 43 où la multimérisation desdits polypeptides est favorisée par une interaction protubérance-dans-cavité entre les domaines de multimérisation.
- 50 45. Méthode de la revendication 43, où chaque domaine variable de chaîne légère est identique.
 - 46. Méthode de la revendication 43, où les premier et second domaines variables de chaîne légère diffèrent l'un de l'autre mais ont au moins 90% d'identité des séquences d'acides aminés.
- 47. Méthode de la revendication 46 où les premier et second domaines variables de chaîne légère ont au moins 95% d'identité des séquences d'acides aminés.
 - 48. Méthode de la revendication 47, où les première et seconde chaînes légères ont au moins 98% d'identité des

séquences.

- 49. Méthode de la revendication 48, où les premier et second domaines variables de chaîne légère ont au moins 99% d'identité des séquences.
- 50. Méthode selon la revendication 1, où l'anticorps multispécifique est sélectionné dans le groupe consistant en antirécepteur ECD leptine humain (Ob-R)/anti-récepteur tyrosine kinase HER3 humain et anti-récepteur tyrosine kinase (Mp1)/thrombopeptine humaine/anti-récepteur tyrosine kinase HER3 humain.
- 51. Anticorps multispécifique de la revendication 17 qui est sélectionné dans le groupe consistant en anti-récepteur 10 ECD leptine humain (Ob-R)/anti-récepteur tyrosine kinase HER3 humain et anti-récepteur tyrosine kinase (Mp1) thrombopoiétine humain/anti-récepteur tyrosine kinase HER3 humain.
 - 52. Composition comprenant l'anticorps multispécifique de la revendication 1 et un support.
 - 53. Cellule hôte de la revendication 28, où l'anticorps multispécifique est sélectionné dans le groupe consistant en anti-récepteur ECD leptine humain (Ob-R)/anti-récepteur tyrosine kinase HER3 humain et anti-récepteur tyrosine kinase thrombopoiétine humain (Mp1)/anti-récepteur tyrosine kinase HER3 humain.

20

15

5

25

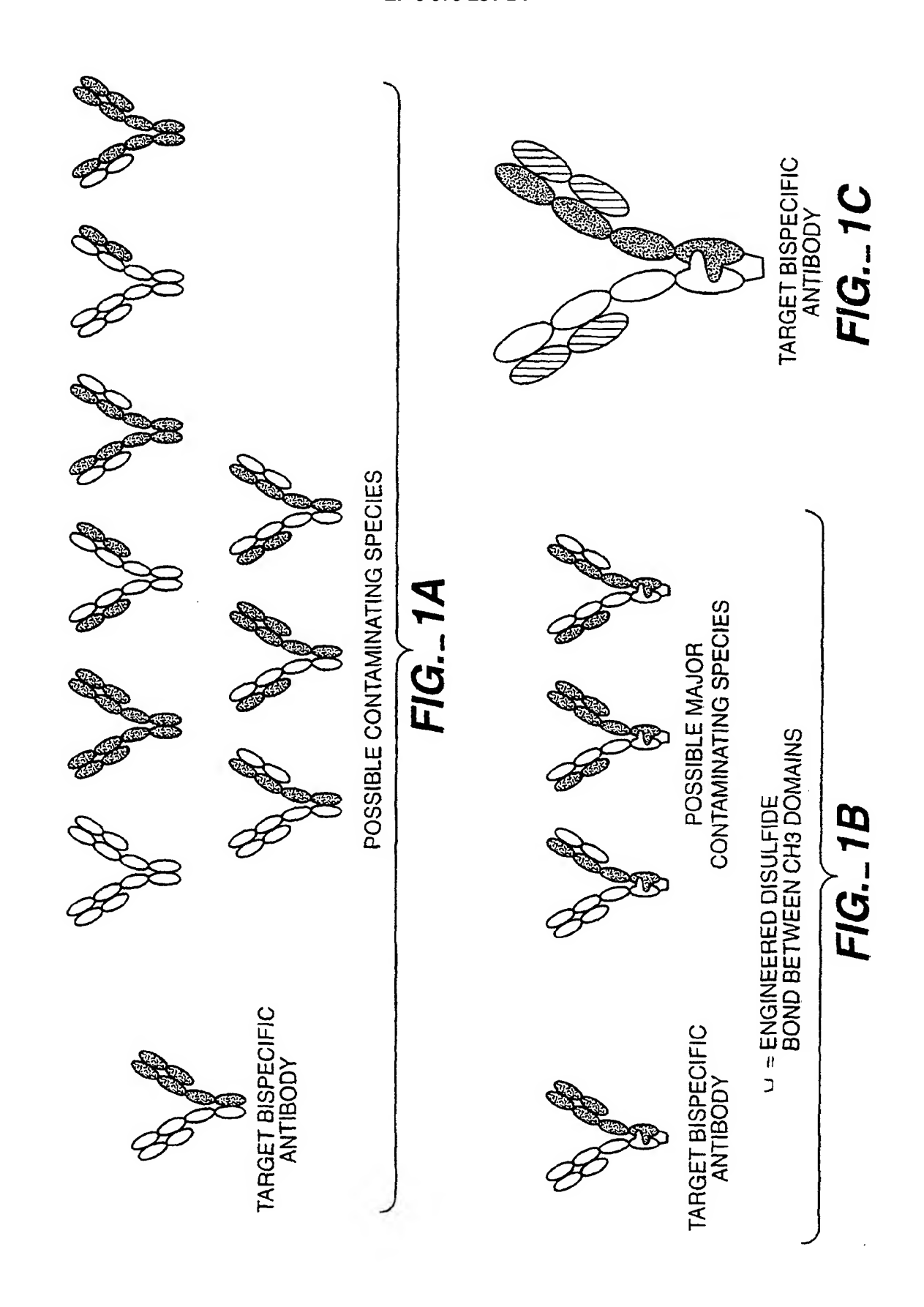
30

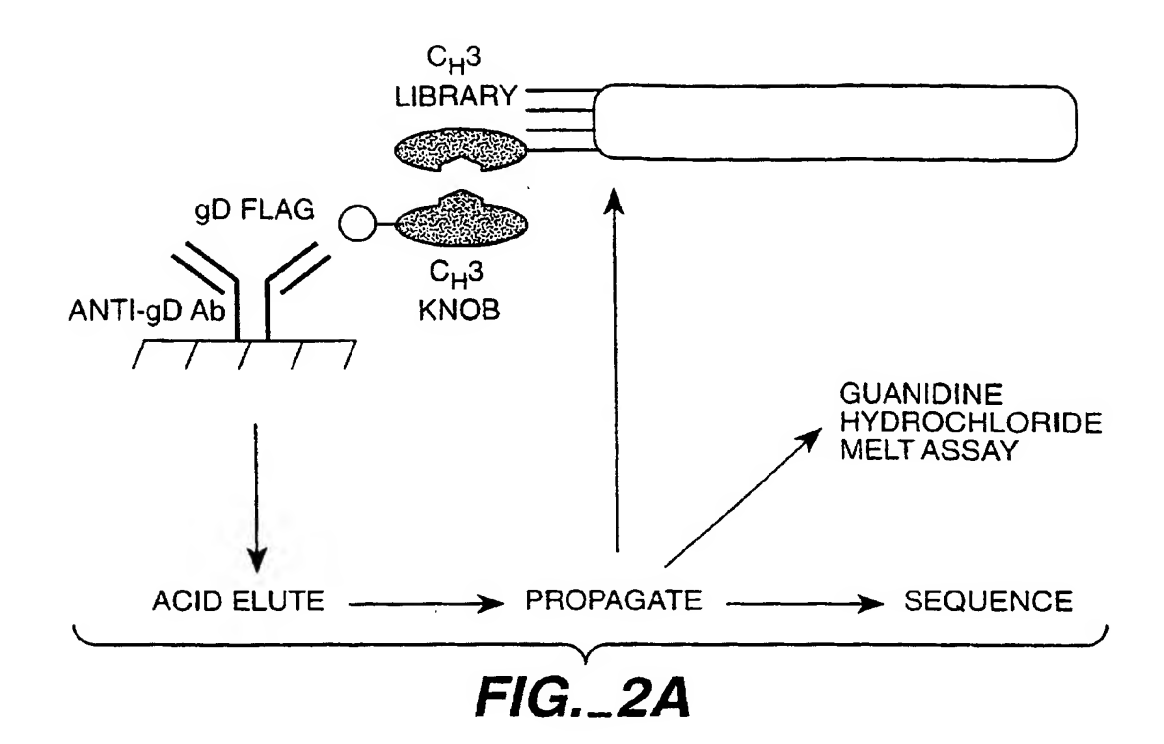
35

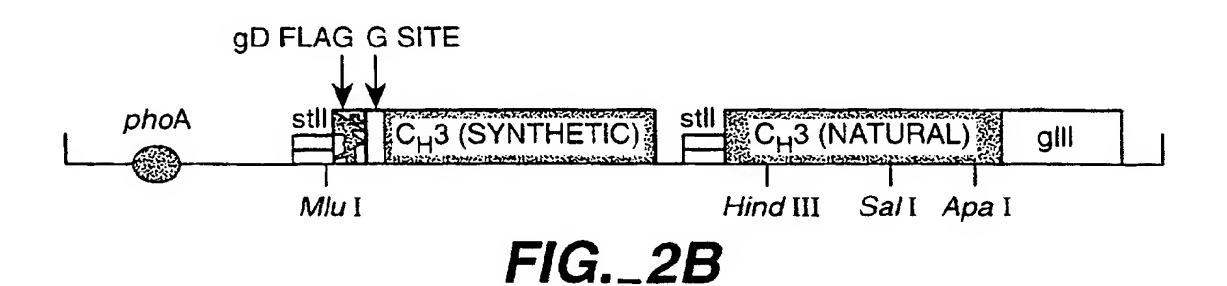
40

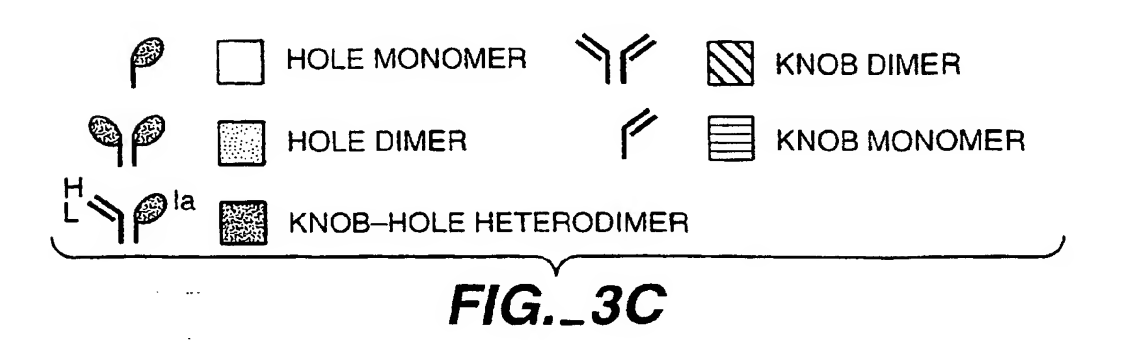
45

50









150 AACGCGTACGCTCTGAAAATGGCGGACCCGAACCGTTTTCGTGGTAAAGATCTGGCTGCACACTACGGCCAGCCG CGGGAACCTCAGGTGTATACCCTGCCACCGTCTCGAGAAAATGACTAAAAACCAGGTCTCTCTGTGGTGCTG 225 300 375 CGTGGAATGGGAAAGCAACGGTCAACCGGAAAAAAACTACAAAACC CTTCTTTCTGTATTCGAAGCTGACCGTTGACAAAAGCCGTTGGCAG CAAGGCAACGTTTTCAGCTGTTATGCACGAGGCCTTGCACAACCACTACACCAGAAAAGCCTGTCCCTG 450 CATCTATGTTCGTTTTTTTTGCTACAACGCGTACGCTGGGCAGCCCCGAGAACCACAGGTGTACACCTGC H CH3 U × Ŋ 350 H <u>۸</u> 366 Ö × 124 × 390 440 H Ч Ø site ď > ß Z × × H Q Ü A > 回 Q Z ρι <u>۸</u> OI > ρ, H K StII 闰 O H Z >1 × K 360 A Ö H 工 **^**1 Σ ρı 幺 H Z Z O^t U E t) ល Ξ Ö PC, 闰 回 **>**+ H K 闰 3 >+ 380 430 DC, [24 闽 4 Ø > Z Ø GTCAAAGGTTTCTATCCGAGCGATATCGC ACTCCACCGGTGCTGGATTGGCTC Σ H ρι А Ö > Н H A 占 S MA gD flag Ω 350 400 H S C Ø Ω Y A L K <u>ዓ</u> P Q Y P P V L D FI V ۸ 1 G F Į#ł 0 > N × Ü Z **स** स 370 420 K D. stII MluI ល > Oi Ø đ

FIG. 2C-1

525

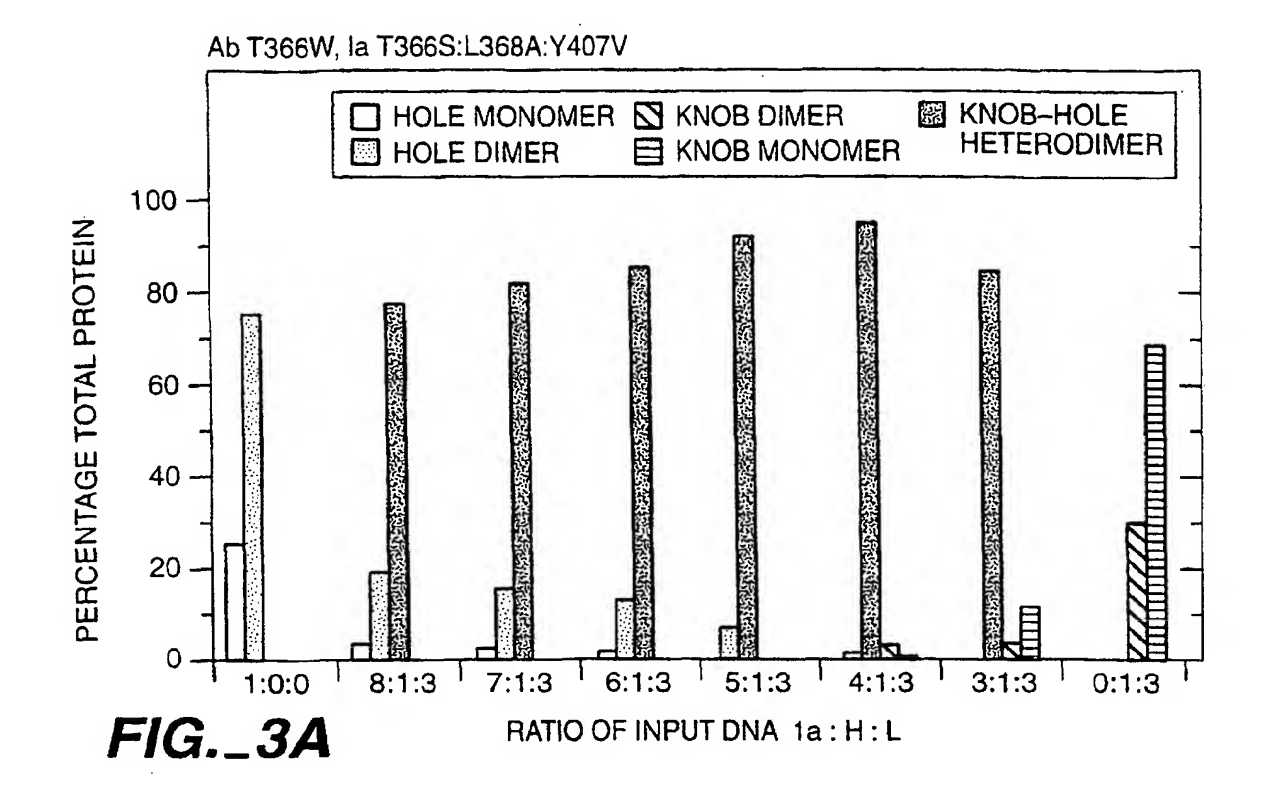
CH3

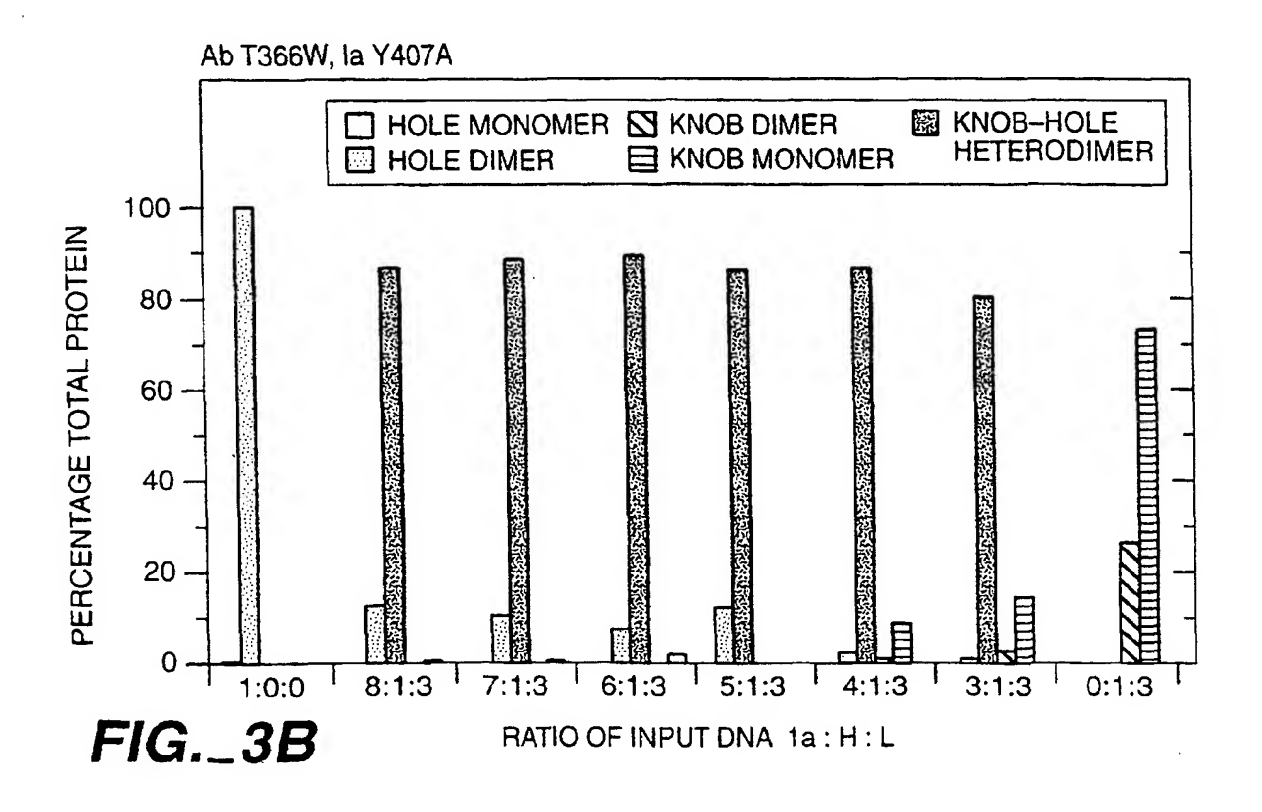
Apar

009 CCCCATCCCGGGAAGATGACCAAGAACCAGGTAAGCTTGTACTGCCTGGTCAAAGGCTTCTATCCCAGCGACA 750 GCTCCTTCTTCTTACAGCTTTCTCGTCGACAAGAGCAGGTGGCAGCAGGGGGAACGTCTTCTTATGCTCCG TCGCCGTGGAGTGGGGAATGGGCAGCCGGAGAACAACTACAAGACCACGCCTCCCGTGCTGGACTCCGACG 400 TGATGCATGAGGCTCTGCACCACTACACGCAGAGAGCCTCTCCCTGTCTCCGGGTAATAGGGGCCC Ŋ Ø Ų щ U Ω S K Am × H Ī4 > > Ω, Z Ü Ü 370 420 × D₄ p4 O H > Ø Ø 368 H H Ø H Ų 3 × Ø 366 × O. **>** H 390 440 H Ø Z HindIII Ø × Z × 国 Ω O) SalI H D4 0 > OI Z H × 360 410 **O** × Ħ Н E [in Z Z Ŋ 耳 Z 田 407 H **>**+ 闰 i I ď 臼 380 430 ĸ 团 > Ħ Ø [±4 X K ф W Н > U

FIG. 2C-2

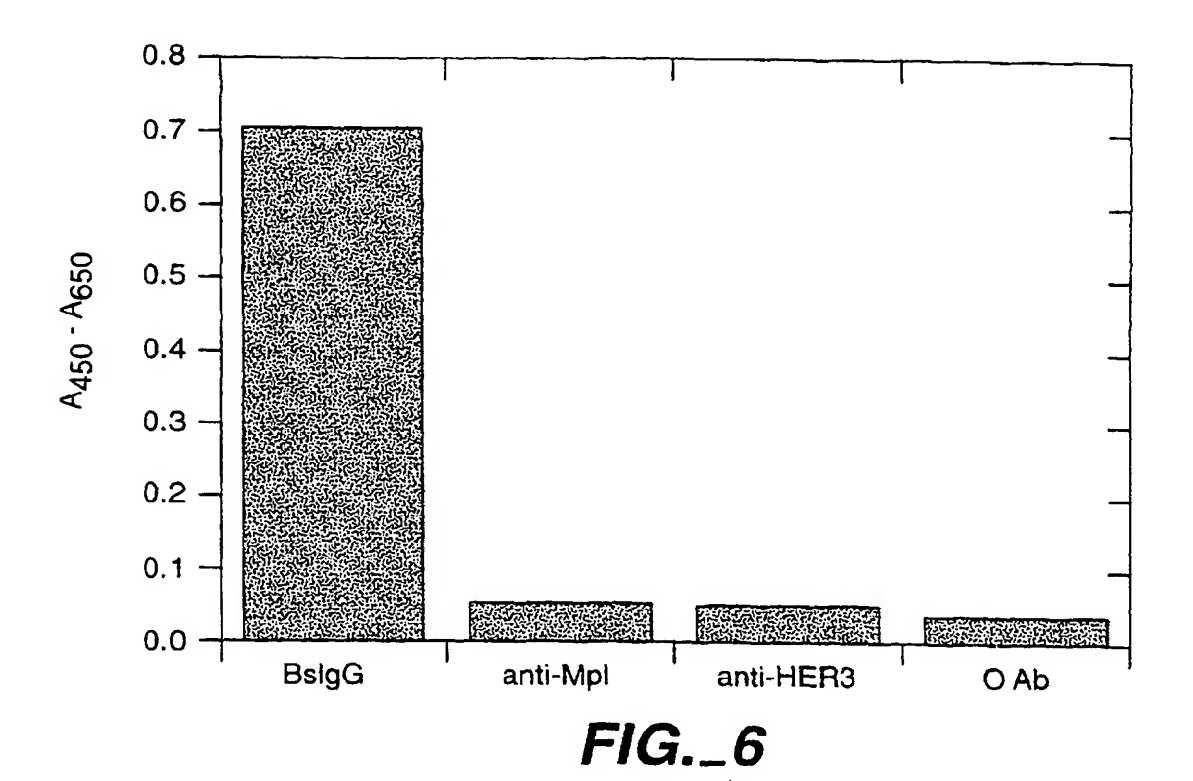
2 FIG. 20-2

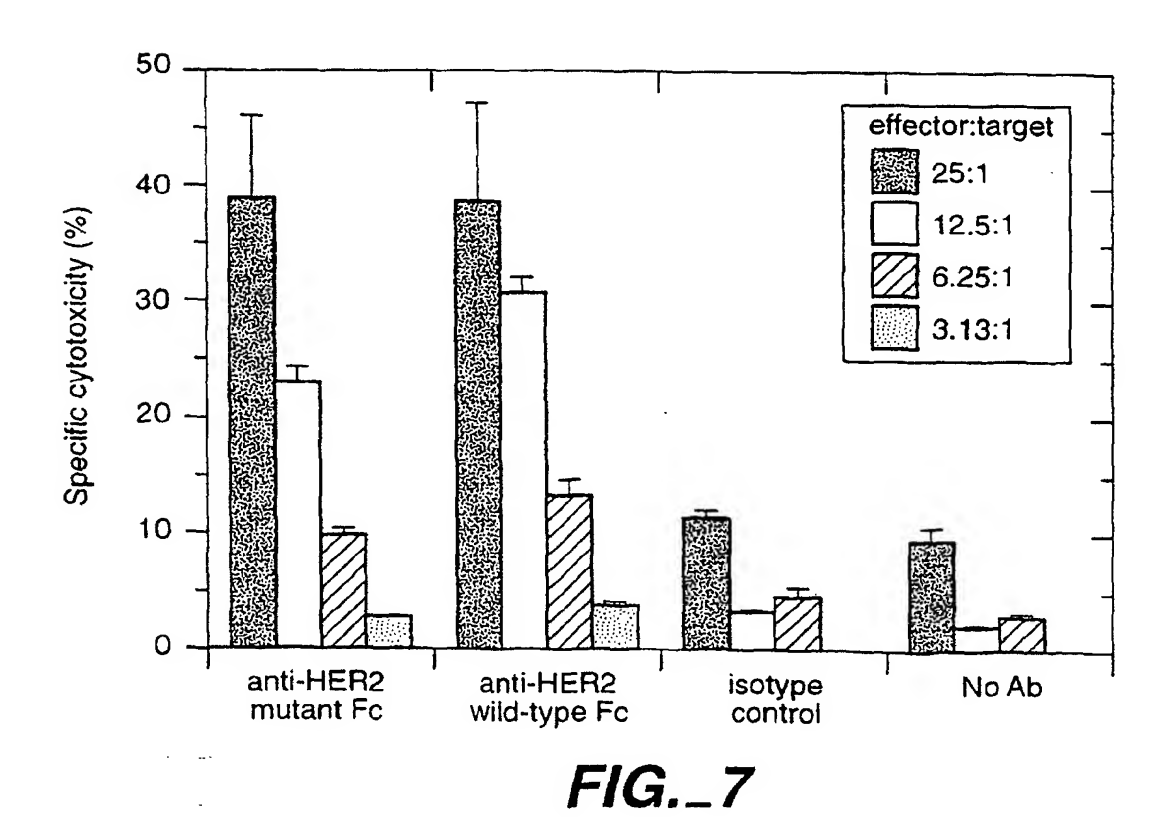




	H	20	abc 30	40	ß	50			
Ax1.78	QSVLTQPASVSGSPGQSITISCTGTSSDVGGYNYVSWYQQHPGKAPKLMIYEGSKRPSGV	PGQSITISCTGT	SSDVGGYNYV	SWYQQHPGKA	PKLMIY	EGSKF	PSG	>	
Rse.23	QSVLTQPASVSGSPGQSITISCTGTSSDVGGYNYVSWYQQHPGKAPKLMIYEGSKRPSGV	egsitiscign	SSDVGGYNYV	SWYQQHPGKA	PKLMIY	EGSKE	BSa	>	
IGER.MAT2C1G11	QSVLTQPASVSGSPGQSITISCTGTSSDVGGYNYVSWYQQHPGKAPKLMIYEGSKRPSGV	GOSITISCIGI	SSDVGGYNYV	SWYQQHPGKA	PKLMIY	EGSKF	(PSG	>	
GCSFR.A4	QSVLTQPASVSGSPGQSITISCTGTSSDVGGYNYVSWYQQHPGKAPKLMIYEGSKRPSGV	COSITISCIG	SSDVGGYNYV	SWYQQHPGKA	PKLMIY	EGSKE	PSG	>	
Rse.04	QSVLTQPASVSGSPGQSITISCTGTSSDVGGYNYVSWYQQHPGKGPKLMIYEGSKRPSGV	GOSITISCTGT	SSDVGGYNYV	SWYQQHPGKG	PKLMIY	EGSKF	PSG	>	
obr.4	QSVLTQPASVSGSPGQSITISCTGTSSDVGGYNYVSWYQQHPGKAPKLMIYEGSKRPSGV	GOSITISCIG	SSDVGGYNYV	SWYQQHPGKA	PKLMIY	EGSKA	(PSG	>	
Rse.20	osvltopasvsgspgositisctgtssdvggynyvswyqqhpgkgpklmiyegskrpsgv	GOSITISCIGE	SSDVGGYNYV	SWYQQHPGKG	PKLMIY	EGSKA	PSG	>	
Rse.15	QSVLTQPASVSGSPGQSITISCTGTSSDVGGYNYVSWYQQHPGKAPKLMIYEGSKRPSGV	GOSITISCTGT	SSDVGGYNYV	SWYQQHPGKA	PKLMIY	EGSKR	PSG	>	
vegf.5	QSVLTQPASVSGSPGQSITISCTGTSSDVGGYNYVSWYQQHPGKAPKLMIYEGSKRPSGV	GOSITISCIGI	SSDVGGYNYV	SWYQQHPGKA	PKLMIY	EGSKR	PSG	>	
		=#=	##########			###			
			CDR L1			COR	17		
	02 29	80	06	a 100					
Ax1.78	SNRFSGSKSGNTASLTISGLQAEDEADYYCSSYTTRSTRVFGGGTKLTVL	LTISGLQAEDE	ADYYCSSYTT	RSTRVFGGGTI	KLTVL	(SEQ	H	NO:	14)
Rse.23	SNRFSGSKSGNTASLTISGLQAEDEADYYCSSYTTRSTRVFGGGTKLTVL	LTISGLQAEDE	ADYYCSSYTT	RSTRVFGGGTI	KLTVL	(SEQ	CI	NO:	15)
IGER. MAT2C1G11	SNRFSGSKSGNTASLTISGLQAEDEADYYCSSYTTRSTRVFGGGTKLTVL	LTISGLQAEDE	ADYYCSSYTT.	RSTRVFGGGT	KLTVL	(SEQ	CI	NO:	16)
GCSFR.A4	SNRFSGSKSGNTASLTISGLQAEDEADYYC	LTISGLQAEDE	ADYYCSSYTT	SSYTTRSTRVFGGGTKLTVL		(SEQ	ID	NO:	17)
Rse. 04	SNRFSGSKSGNTASLTISGLQAEDEADYYCSSYTTRSTRVFGGGTKLTVL	LTISGLQAEDE	ADYYCSSYTT	RSTRVFGGGT		(SEQ	ID	NO:	18)
obr.4	SNRFSGSKSGSTASLTISGLQAEDEADYYCSSYTTRSTRVFGGGTKLTVL	LTISGLQAEDE	ADYYCSSYTTI	RSTRVFGGGTF		ÕES)	ID	NO:	19)
Rse.20	SNRFSGSKKGNTASLTISGLQAEDEADYYCSSYTTRSTRVFGGGTKLTVL	LTISGLQAEDE	ADYYCSSYTT	RSTRVFGGGTR		(SEQ	QI	NO:	20)
Rse.15	SNRFSGSKSGNTASLTISGLQAEDEADYYCSSYTTRSTRVFGGGTKLTVL	LTISGLQAEDE	ADYYCSSYTTI	rstrvfgggtr		ÕES)	ID	NO:	21)
vegf.5	SNRFSGSKSGNTASLTISGLQAEDEADYYCSSYTTRSTRVFGGGTKLTVL	LTISGLQAEDE	ADYYCSSYTT	RSTRVFGGGTR		(SEQ	ID	NO:	22)
			###	######					
			CDR	L3					
		1	1						

24) (SEQ ID NO: ID NO: (SEQ ADSVKGRFTI SRDNSKNTLYLQMNRLRAEDTAVYYCARDNGWELTDWYFDLWGRGTMVTVSS OVOLVOSGGGLVQPGGSLRLSCAASGFTFSSYEMN--WVRQAPGKGLEWVSGISGSGGSTYY NPSLKSRVTI SVDTSKNQFSLKLSSVTAADTAVYYCARVDLEDYGSGASDYWGQGTLVTVSS EVQLVESGPGLVKPSQTLSLTCTVSGGSISSGGYYWSWIRQHPGKGLEWIGYIY-YSGSTYY **** DIOMTOSPSTLSASIGDRVTITCRASEGIYHWLAWYQQKPGKAPKLLIYKASSLASGAPSRF 25) 9 CDR H2 110 (SEQ ID NO: * CDR L2 50 50 100abcde CDR H3 SGSGSGTDFTLTISSLQPDDFATYYCOOYSNYPLTFGGGTKLEIK 40 **** 100 ap 06 CDR H1 CDR L3 CDR L1 30 30 90 apc 80 * 20 20 80 *** * *** * *** 70 10 * 70 her3.18 her3.18 CDR H2 *** * obr. 26 obr.26 9





	H1	H2	НЗ	H4	H5	Н6	H7	Н8	Н9	H10	H11
01	49	47	51	81	60	48	76	51	100	62	51
02	84	79	88	50	48	99	48	88	48	45	88
03	83	82	85	51	50	95	49	85	49	46	85
04	47	50	51	83	77	48	65	51	73	64	51
05	49	47	51	81	60	48	76	51	100	62	51
06	83	79	86	50	50	99	47	86	48	45	86
07	81	100	86	51	49	80	48	86	47	44	86
08	81	100	86	51	49	80	48	86	47	44	86
09	81	100	86	51	49	80	48	86	47	44	86
010	83	79	85	50	49	98	46	85	48	45	85
011	83	80	87	50	49	99	47	87	48	45	87
012	81	100	86	51	49	80	48	86	47	44	86
013	49	47	51	81	60	48	76	51	100	62	51
014	50	50	54	95	67	49	76	54	75	62	54
015	82	79	85	49	48	97	46	85	47	44	85
016	84	80	87	50	49	100	47	87	48	45	87
017	45	44	47	65	62	45	62	47	62	100	47
O18	50	51	50	75	79	50	63	50	66	62	50

01-018: ANTI-Ob-R ANTIBODY CLONES obr. 1, 11, 12, 14, 15, 16, 17, 18, 19, 2, 20, 21, 22, 23, 24, 26, 3, 4, RESPECTIVELY.

H1-H11: ANTI-HER3 ANTIBODY CLONES her3. 1, 3.10, 3.11, 3.12, 3.16, 3.18, 3.19, 3.22, 3.3, 3.4, 3.7, RESPECTIVELY.

FIG._8